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- Key terms

E PHOSPHOLIPASE D/CN 5 L1 154 S PHOSPHOLIPASE D ?/CN

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FILE COVERS 1907 - 3 May 2006 VOL 144 ISS 19 FILE LAST UPDATED: 2 May 2006 (20060502/ED)

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http://www.cas.org/infopolicy.html

L1
154 SEA FILE=REGISTRY ABB=ON PLU=ON PHOSPHOLIPASE D ?/CN
L2
4852 SEA FILE=CAPLUS ABB=ON PLU=ON L1 OR (PHOSPHOLIPASE OR
PHOSPHO LIPASE OR LECITHINASE) (1W) D OR (PHOSPHATIDYLCHOLINE
OR PHOSPHATIDYL CHOLINE) (W) (PHOSPHOHYDROLASE OR PHOSPHO
HYDROLASE)

L4 8 SEA FILE=CAPLUS ABB=ON PLU=ON L2 AND ?NEISSER?

L4 ANSWER 1 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN

ED Entered STN: 17 Dec 2004

ACCESSION NUMBER: 2004:1080507 CAPLUS

DOCUMENT NUMBER: 142:54745

TITLE: Vaccine and compositions comprising a

neisserial phospholipase

D for the prevention and treatment of

neisserial infections

INVENTOR(S): Apicella, Michael A.; Edwards, Jennifer L.

PATENT ASSIGNEE(S): USA

SOURCE: U.S. Pat. Appl. Publ., 103 pp., Cont.-in-part of

U.S. Ser. No. 621,184.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.					DATE		APPLICATION NO.						DATE			
US 2004253222				A1 200 A1 200		2004 2003	20041216 20030529		US 2003-665990 US 2002-66551					20020131		
WO																
	W:	CH,	CN,	co,	CR,	CU,	AU, CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,
							HR,									
							LS,									
							NZ,									
		-	-				ТJ,	TM,	TN,	TK,	TT,	TZ,	UA,	uG,	UZ,	VC,
	DE7.	•	•	ZA,	•		Mari	М7	N 7 7	CD.	CT	C 7	m o	IIC	7 M	77.547
	RW:						MD,									
							FR,									
							TR,									
							TD,		ъо,	CF,	CG,	CI,	CP1,	GA,	GIV,	GQ,
PRIORITY	APP				NE,	JN,	10,	10	1	US 2	001-	2660	70P		P 2	0010131
									1	US 2	001-	3103	56P		P 2	0010806
									1	US 2	001-	3444	52P		P 2	0011023
				-					1	US 2	002-	6655	1		A2 2	0020131
									1	US 2	003-	6211	84		A2 2	0030715
									1	US 2	003-	6659	90		A2 2	0030919

The present invention provides a polypeptide, polynucleotide, vaccine, AB and a method of vaccination effective to immunize a mammal against a neisserial infection, e.g., an infection caused by Neisseria gonorrhoeae or Neisseria meningitidis by using a neisserial phospholipase D (PLD) polypeptide in combination with a physiol.-acceptable, non-toxic vehicle. In addition, the invention provides a transgenic Neisseria bacterium comprising a disrupted pld gene wherein the bacterium has reduced phospholipase D activity as compared to the phospholipase D activity of a corresponding wild-type Neisseria. IT 808201-07-2P, Phospholipase D ( Neisseria gonorrhoeae) 808201-30-1P 808201-31-2P 808201-32-3P RL: BPN (Biosynthetic preparation); BSU (Biological study, unclassified); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (amino acid sequence; vaccine and compns. comprising neisserial phospholipase D for the prevention and treatment of neisserial infections) ANSWER 2 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN L4Entered STN: 07 Nov 2003 ACCESSION NUMBER: 2003:873418 CAPLUS 139:379737 DOCUMENT NUMBER: TITLE: Gonococcal phospholipase D modulates the expression and function of complement receptor 3 in primary cervical epithelial cells Edwards, Jennifer L.; Entz, David D.; Apicella, AUTHOR(S): Michael A. Department of Microbiology, University of Iowa, CORPORATE SOURCE: Iowa City, IA, 52242, USA Infection and Immunity (2003), 71(11), 6381-6391 SOURCE: CODEN: INFIBR; ISSN: 0019-9567 PUBLISHER: American Society for Microbiology DOCUMENT TYPE: Journal LANGUAGE: English CR3-mediated endocytosis is a primary mechanism by which Neisseria gonorrhoeae elicits membrane ruffling and cellular invasion of the cervical epithelia. The authors' data indicate that, upon infection of cervical epithelia, N. gonorrhoeae specifically releases proteins, including a phospholipase D (PLD) homolog, which facilitate membrane ruffling. To elucidate the function of gonococcal PLD in infection of the cervical epithelia, the authors constructed an N. gonorrhoeae PLD mutant. By comparative association and/or invasion assays, the authors demonstrated that PLD mutant gonococci are impaired in their ability to adhere to and to invade primary cervical cells. This defect can be rescued by the addition of supernatants obtained from wild-type-infected cell monolayers but not by exogenously added Streptomyces PLD. The decreased level of total cell association (i.e., adherence and invasion) observed for mutant gonococci is, in part, attributed to the inability of these bacteria to recruit CR3 to the cervical cell surface with extended infection. Using electron microscopy, the authors demonstrate that gonococcal PLD may be necessary to potentiate membrane ruffling and clustering of gonococci on the cervical cell surface. These data may be indicative

Searcher : Shears 571-272-2528

of the inability of PLD mutant gonococci to recruit CR3 to the cervical cell surface. Alternatively, in the absence of gonococcal

PLD, signal transduction events required for CR3 clustering may not be activated. Collectively, the authors' data indicate that PLD augments CR3-mediated gonococcus invasion of and survival within cervical epithelia.

REFERENCE COUNT:

THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 3 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN

ED Entered STN: 11 Apr 2003

ACCESSION NUMBER: 2003:282761 CAPLUS

DOCUMENT NUMBER: 138:300147

TITLE: Sensitive and rapid detection of pathogenic

organisms and toxins using fluorescent polymeric

lipids

INVENTOR(S): Moronne, Mario Manuel; Charych, Deborah H.; Nagy,

Jon O.

PATENT ASSIGNEE(S): Regents of the University of California, USA

SOURCE: PCT Int. Appl., 91 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE: Eng FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

1	PATENT NO.			KIND DATE		APPLICATION NO.					DATE						
	WO 2003029479 WO 2003029479										20020809			09			
	W		AG, CO,	•	•	•	•	•	•	-		_	-	-	-		
			GH,	•	•	•	•	•	•	•							
			), NZ, I, TN,	•	•	•	•	•	•	•	•	-	-	•	-	TJ,	
	P		, GM, , KG,	•	•	•		•		-	-			-	-	-	
		В	E, ES, BJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	
t	US 20	03129	618		A1		2003	0710	1	US 2	002-	2157	36		2	00208	09
1		: A7	', BE, ', IE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	09
PRIOR	ITY A				±1± /	, DV	,	110,								00108	10
					·				,	us 2	002-	2157	36	i	A 2	00208	09
									1	WO 2	002-	US25	486	1	₩ 2	00208	09

- AB The present invention relates to methods and compns. for the detection of analytes using the fluorescence that occurs in polymeric material in response to selective binding of analytes to the polymeric materials. In particular, the present invention allows for the fluorescent detection of membrane modifying reactions and analytes responsible for such modifications and for the screening of reaction inhibitors.
- L4 ANSWER 4 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN

Entered STN: 24 Oct 2001 ED

2001:772087 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 135:341173

Nucleic acid-coupled colorimetric analyte TITLE:

detectors using self-assembling polydiacetylene

liposomes

INVENTOR(S): Charych, Deborah H.; Jonas, Ulrich

PATENT ASSIGNEE(S): Regents of the University of California, USA U.S., 96 pp., Cont.-in-part of U.S. Ser. No. SOURCE:

461,509.

CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 11

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
US 6306598 US 6001556 EP 1460423 R: AT, BE, CH, PT, IE	B1 20011023 A 19991214 A1 20040922		19990621 19960126 19960213 SE, MC,
US 6183772 US 6022748 US 6080423 US 6180135 US 6468759 CA 2330937 JP 2004500006 US 6395561 US 6485987 US 2001026915	B1 20010206 A 20000208 A 20000627 B1 20010130 B1 20021022 AA 19991229 T2 20040108 B1 20020528 B1 20021126 A1 20011004		19960301 19970829 19971006 19971006 19980302 19990622 19990622 19991214 20000208 20001211
US 6660484 PRIORITY APPLN. INFO.:	B2 20031209	US 1992-976697	A2 19921113
		US 1993-159927	A2 19931130
		US 1994-289384	B2 19940811
		US 1994-289384	B2 19940811
•		US 1994-328237	B2 19941024
		US 1995-389475	B3 19950213
		US 1995-389475	B2 19950213
		US 1996-592724	A3 19960126
		US 1996-609312	A2 19960301
		US 1997-38383P	P 19970214
		US 1997-39749P	P 19970303
		US 1997-50496P	P 19970623
		US 1997-920501	A3 19970829

571-272-2528 Searcher Shears

US	1997-944323	A2	19971006
US	1998-23898	A2	19980213
US	1998-33557	A2	19980302
US	1998-90266P	P	19980622
US	1998-103344	A2	19980623
US	1999-461509	A2	19991214
US	2000-500295	A2	20000208
US	1992-982189	В2	19921125
EP	1996-906444	АЗ	19960213
US	1997-944257	АЗ	19971006
US	1999-337973	Α	19990621
WO	1999-US14029	W	19990622
US	1999-170190P	P	19991210

AB The present invention relates to methods and compns. for the direct detection of analytes and membrane conformational changes through the detection of color changes in biopolymeric materials. In particular, the present invention provides for the direct colorimetric detection of analytes using nucleic acid ligands at surfaces of polydiacetylene liposomes and related mol. layer systems. Liposomes were prepared from a lipid mixture of 95% 5,7-docsoadiynoic acid and 5% 5,7-docsoadiynoate succinimide. The liposome solution was photopolymd. with UV light (254 nm) and then reacted with RGGGAATTCGTR (R = OP(OH)(O)OCH2(CH2OH)CH(CH2)4NH2) to make a probe.

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 5 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN L4

Entered STN: 19 Apr 2000 ED

2000:250828 CAPLUS ACCESSION NUMBER:

132:261300 DOCUMENT NUMBER:

Complete DNA sequence of a serogroup A strain of TITLE:

Neisseria meningitidis 22491

AUTHOR(S): Parkhill, J.; Achtman, M.; James, K. D.; Bentley,

S. D.; Churcher, C.; Klee, S. R.; Morelli, G.; Basham, D.; Brown, D.; Chillingworth, T.; Davies,

R. M.; Davis, P.; Devlin, K.; Feltwell, T.;

Hamlin, N.; Holroyd, S.; Jagels, K.; Leather, S.; Moule, S.; Mungall, K.; Quail, M. A.; Rajandream, M.-A.; Rutherford, K. M.; Simmonds, M.; Skelton, J.; Whitehead, S.; Spratt, B. G.; Barrell, B. G.

The Sanger Centre, The Wellcome Trust Genome CORPORATE SOURCE:

Campus, Hinxton, Cambridge, CB10 ISA, UK

Nature (London) (2000), 404(6777), 502-506 SOURCE:

CODEN: NATUAS; ISSN: 0028-0836

PUBLISHER: Nature Publishing Group

DOCUMENT TYPE: Journal LANGUAGE: English

AB The complete genome sequence was determined for a serogroup A strain of Neisseria meningitidis, Z2491. The sequence is 2,184,406 bp in length, with an overall G+C content of 51.8%, and contains 2121 predicted coding sequences. The most notable feature of the genome is the presence of many hundreds of repetitive elements, ranging from short repeats, positioned either singly or in large multiple arrays, to insertion sequences and gene duplications of one kilobase or more. Many of these repeats appear to be involved in genome fluidity and antigenic variation in this important human pathogen.

IT 263000-67-5

RL: BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study)

(amino acid sequence; complete DNA sequence of a serogroup A strain of Neisseria meningitidis Z2491)

REFERENCE COUNT:

THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 6 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN

30

ED Entered STN: 30 Dec 1999

ACCESSION NUMBER: 1999:819529 CAPLUS

DOCUMENT NUMBER: 132:60102

TITLE: Nucleic acid-coupled colorimetric analyte

detectors using self-assembling polydiacetylenic

materials

Patent

INVENTOR(S): Charych, Deborah H.; Jonas, Ulrich

PATENT ASSIGNEE(S): Regents of the University of California, USA

SOURCE: PCT Int. Appl., 176 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 11

PATENT INFORMATION:

PA.	PATENT NO.				KIND DATE		APPLICATION NO.					DATE					
WO	WO 9967423			A1		1999	1229		WO 1	1999-	US14	029			19990	0622	
	W:	AU,	CA,	JP													
	RW:	AT,	BE,	CH,	CY,	DE,	DK,	ES,	FI,	FR,	, GB,	GR,	ΙE,	ΙT,	LU	MC,	,
		NL,	PT,	SE													
CA	2330	937			AA		1999	1229		CA :	1999-	2330	937			19990	0622
AU	9947	047		_	A1		2000	0110		AU 1	1999-	4704	7		:	L9990	0622
AU	7486	44			В2		2002	0606									
EP	1112	377			A1		2001	0704		EP 3	1999-	9305	22			19990	0622
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	, IT,	LI,	LU,	NL,	SE	MC,	,
		PT,	ΙE,	FΙ													
JP	2004	5000	06		Т2		2004	0108			2000-					L9990	
PRIORIT	Y APP	LN.	INFO	. :						US :	1998-	9026	6P		P :	19980	0622
										מון	1999-	3379	73		Δ.	19991	1621
										05 .	1000	5515	, ,		4.		, , ,
										WO :	1999-	US14	029		W :	19990	0622

AB The present invention relates to methods and compns. for the direct detection of analytes and membrane conformational changes through the detection of color changes in biopolymeric materials. In particular,

the present invention provides for the direct colorimetric detection of analytes using nucleic acid ligands at surfaces or polydiacetylene liposomes and related mol. layer systems. Synthetic schemes are provided for the preparation and immobilization of polydiacetylenic materials with various head groups.

REFERENCE COUNT:

AUTHOR(S):

THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 7 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN

9

ED Entered STN: 02 Apr 1994

ACCESSION NUMBER: 1994:158350 CAPLUS

DOCUMENT NUMBER: 120:158350

TITLE: Involvement of phospholipid end groups of group C

Neisseria meningitidis and Haemophilus

influenzae type b polysaccharides in association with isolated outer membranes and in immunoassays Arakere, Gayathri; Lee, Ann L.; Frasch, Carl E.

CORPORATE SOURCE: Cent. Biol. Eval. Res., Div. Bacterial Prod.,

Bethesda, MD, 20892, USA

SOURCE: Journal of Bacteriology (1994), 176(3), 691-5

CODEN: JOBAAY; ISSN: 0021-9193

DOCUMENT TYPE: Journal LANGUAGE: English

There are several bacterial polysaccharides (PSs) which contain a AB terminal lipid moiety. It has been postulated that these terminal lipid mols. anchor the PSs to the outer membrane of the bacteria. The authors show here that incubation of native PS from group C Neisseria meningitidis or Haemophilus influenza type b with isolated outer membrane vesicles results in association of a portion of the PS with the vesicles. Removal of the terminal lipid from the PS by treatment with phospholipase A2 or phospholipase p eliminates this association In other studies, it was shown that delipidated PSs are not suitable as solid-phase antigens in a currently used ELISA. Measurement of antibody units in the reference sera by using delipidated PSs as antigens in an ELISA yielded negligible absorbance compared with native PSs when methylated human serum albumin was used to coat the PSs to the plate. Nevertheless, phospholipase A2 and phospholipase D treatment did not noticeably affect antigenic epitopes, since soluble group C PS without the terminal lipid bound antibody as effectively as the native PS did, as measured by a competitive inhibition assay. Both hydrophobic and electrostatic interactions are important for the binding of group C N. meningitidis PS to the ELISA plate, while charge interactions seem to be sufficient for binding the more neg. charged H. influenzae type b PS.

L4 ANSWER 8 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN

ED Entered STN: 01 May 1993

ACCESSION NUMBER: 1993:167635 CAPLUS

DOCUMENT NUMBER: 118:167635

TITLE: Process for converting bacterial lipid-containing

capsular polysaccharide into lipid-free

polysaccharide

INVENTOR(S): Lee, Ann L.; Sitrin, Robert D.; Manger, Walter E.;

Rienstra, Mark S.

PATENT ASSIGNEE(S): Merck and Co., Inc., USA SOURCE: Eur. Pat. Appl., 28 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	TENT NO.					API	PLICATION NO.		D	ATE
EP	528635			A1		EP	1992-307395		1	9920812
	R: AT,	BE,	CH,	DE,	DK, ES, FR,	GB, GI	R, IE, IT, LI	, LU,	NL,	PT, SE
	5314811	•	•	A	19940524	US	1992-909346		1	9920713
WO	9304183			A1	19930304	_ WO	1992-US6301		1	9920729
	W: BG,	CS,	FI,	HU,	NO, PL, RO,	RÚ				
							1992-2075681		1	9920810
CA	2075681			С	20030325					
AT	176929			E	19990315	AT	1992-307395		1	9920812
AU	9221054			A1	19930218	AU	1992-21054		1	9920814
ZA	9206131			Α	19930428	ZA	1992-6131		1	9920814
JP	05209002						1992-217011		1	9920814
CN	1071699			Α	19930505	CN	1992-110465		1	9920815
NO	9400519			Α	19940215	NO	1994-519		1	9940215
PRIORITY	Y APPLN.	INFO	. :			US	1991-746523	1	A 1	9910816
						US	1992-909346	1	A 1	9920713
						WO	1992-US6301	1	A 1	9920729

AB A process for converting lipid-containing bacterial capsular polysaccharide, such as lipo-polyribosyl ribitol phosphate (lipo-PRP), into lipid-free, endotoxin-free polysaccharide, such as PRP, is claimed. The process comprises solubilizing a polysaccharide-containing powder derived from the bacterial culture, cleaving the covalently bound lipid from the polysaccharide, and removing the lipids and endotoxin. Thus, a phenol-inactivated pre-phenol PRP powder derived from Haemophilus influenzae type b was digested with phospholipase D and the enzyme was removed by phenol extraction After removal of LPS antigen by HP2O chromatog., the lipid-free PRP was prepared by diafiltration and EtOH precipitation The PRP prepared by this

process and by the prior art selective alc. fractionation process were indistinguishable in physicochem. and (in vitro and in vivo) immunogenicity assays.

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FILE 'JICST-EPLUS' ENTERED AT 10:46:11 ON 03 MAY 2006 COPYRIGHT (C) 2006 Japan Science and Technology Agency (JST)

FILE 'JAPIO' ENTERED AT 10:46:11 ON 03 MAY 2006 COPYRIGHT (C) 2006 Japanese Patent Office (JPO) - JAPIO

L5 11 S L4

L6 5 DUP REM L5 (6 DUPLICATES REMOVED)

L6 ANSWER 1 OF 5 WPIDS COPYRIGHT 2006 THE THOMSON CORP on STN

ACCESSION NUMBER:

2005-123122 [13] WPIDS

CROSS REFERENCE: .

2002-619227 [66]

DOC. NO. CPI:

C2005-040896

TITLE:

New transgenic Neisseria bacterium

comprising a disrupted pld gene and a reduced

phospholipase D activity, useful
for preventing or treating neisserial

infections, such as gonorrhea.

DERWENT CLASS:

B04 D16

INVENTOR(S):
PATENT ASSIGNEE(S):

APICELLA, M A; EDWARDS, J L (IOWA) UNIV IOWA RES FOUND

COUNTRY COUNT:

107

PATENT INFORMATION:

PATENT NO	KIND DAT	re week	LA	PG
				· <b>-</b>
FTO 2005010026	31 2005	3002 /200E121+	EM 163	,

WO 2005010036 A1 20050203 (200513) \* EN 163

RW: AT BE BG BW CH CY CZ DE DK EA EE ES FI FR GB GH GM GR HU IE IT KE LS LU MC MW MZ NA NL OA PL PT RO SD SE SI SK SL SZ TR TZ UG ZM ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BW BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE EG ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NA NI NO NZ OM PG PH PL PT RO RU SC SD SE SG SK SL SY TJ TM TN TR TT TZ UA UG UZ VC VN YU ZA ZM ZW

#### APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
WO 2005010036		WO 2004-US22708	20040715

PRIORITY APPLN. INFO: US 2003-665990 20030919; US

2003-621184

20030715

AN 2005-123122 [13] WPIDS

CR 2002-619227 [66]

AB W02005010036 A UPAB: 20050224

NOVELTY - A transgenic Neisseria bacterium comprising a disrupted pld gene, is new. The bacterium has reduced phospholipase D (PLD) activity as compared to the phospholipase D activity of a corresponding wild-type Neisseria.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated and purified polynucleotide encoding a PLD from a Neisseria bacterium;
  - (2) an isolated and purified polypeptide that is encoded by the

above polynucleotide and that comprises phospholipase p from a Neisseria bacterium;

- (3) a vaccine comprising an immunogenic amount of a PLD polypeptide from Neisseria, which amount immunizes a patient against a neisserial infection, in combination with a physiological, non-toxic vehicle;
- (4) protecting a patient against **Neisseria** colonization or infection, comprising administering to the patient an amount of the vaccine mentioned above; and
- (5) preventing infection or colonization of Neisseria in a patient by administering to the patient a compound that inhibits neisserial phospholipase D.

ACTIVITY - Antibacterial; Gynecological.

No biological data given.

MECHANISM OF ACTION - Vaccine.

USE - The composition and methods are useful for preventing or treating neisserial infections, such as gonorrhea. Dwg.0/23

L6 ANSWER 2 OF 5 MEDLINE on STN DUPLICATE 1

ACCESSION NUMBER: 2003496542 MEDLINE DOCUMENT NUMBER: PubMed ID: 14573659

TITLE: Gonococcal phospholipase d

modulates the expression and function of complement receptor 3 in primary cervical epithelial cells.

AUTHOR: Edwards Jennifer L; Entz David D; Apicella Michael A CORPORATE SOURCE: Department of Microbiology, University of Iowa, Iowa

City, Iowa 52242, USA.

CONTRACT NUMBER: 5- 32-AI07343-14T (NIAID)

AI38515 (NIAID) AI45728 (NIAID)

SOURCE: Infection and immunity, (2003 Nov) Vol. 71, No. 11, pp.

6381-91.

Journal code: 0246127. ISSN: 0019-9567.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200311

ENTRY DATE: Entered STN: 24 Oct 2003

Last Updated on STN: 19 Dec 2003 Entered Medline: 20 Nov 2003

AB CR3-mediated endocytosis is a primary mechanism by which Neisseria gonorrhoeae elicits membrane ruffling and cellular invasion of the cervical epithelia. Our data indicate that, upon infection of cervical epithelia, N. gonorrhoeae specifically releases proteins, including a phospholipase D (PLD) homolog, which facilitate membrane ruffling. To elucidate the function of gonococcal PLD in infection of the cervical epithelia, we constructed an N. gonorrhoeae PLD mutant. By comparative association and/or invasion assays, we demonstrated that PLD mutant gonococci are impaired in their ability to adhere to and to invade primary cervical cells. This defect can be rescued by the addition of supernatants obtained from wild-type-infected cell monolayers but not by exogenously added Streptomyces PLD. The decreased level of total cell association (i.e., adherence and invasion) observed for mutant gonococci is, in part, attributed to the inability of these bacteria to recruit CR3 to the cervical cell surface with extended infection. Using electron microscopy, we demonstrate that gonococcal PLD may be

necessary to potentiate membrane ruffling and clustering of gonococci on the cervical cell surface. These data may be indicative of the inability of PLD mutant gonococci to recruit CR3 to the cervical cell surface. Alternatively, in the absence of gonococcal PLD, signal transduction events required for CR3 clustering may not be activated. Collectively, our data indicate that PLD augments CR3-mediated gonococcus invasion of and survival within cervical epithelia.

L6 ANSWER 3 OF 5 MEDLINE on STN DUPLICATE 2

ACCESSION NUMBER: 94131948 MEDLINE DOCUMENT NUMBER: PubMed ID: 8300524

TITLE: Involvement of phospholipid end groups of group C

Neisseria meningitidis and Haemophilus

influenzae type b polysaccharides in association with

isolated outer membranes and in immunoassays.

AUTHOR: Arakere G; Lee A L; Frasch C E

CORPORATE SOURCE: Center for Biologics Evaluation and Research, Division

of Bacterial Products, Bethesda, Maryland 20892.

SOURCE: Journal of bacteriology, (1994 Feb) Vol. 176, No. 3,

pp. 691-5.

Journal code: 2985120R. ISSN: 0021-9193.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199403

ENTRY DATE: Entered STN: 18 Mar 1994

Last Updated on STN: 18 Mar 1994

Entered Medline: 8 Mar 1994

AΒ There are several bacterial polysaccharides (PSs) which contain a terminal lipid moiety. It has been postulated that these terminal lipid moieties anchor the PSs to the outer membrane of the bacteria. Our studies have shown that incubation of native PS from group C Neisseria meningitidis or Haemophilus influenzae type b with isolated outer membrane vesicles results in association of a portion of the PS with the vesicles. Removal of the terminal lipid from the PS by treatment with phospholipase A2 or phospholipase D eliminates this association. In other studies, it was shown that delipidated PSs are not suitable as solid-phase antigens in a currently used enzyme-linked immunosorbent assay (ELISA). Measurement of antibody units in the reference sera by using delipidated PSs as antigens in an ELISA yielded negligible absorbance compared with native PSs when methylated human serum albumin was used to coat the PSs to the plate. Nevertheless, phospholipase A2 and phospholipase D treatment did not noticeably affect antigenic epitopes, since soluble group C PS without the terminal lipid bound antibody as effectively as the native PS did, as measured by a competitive inhibition assay. Both hydrophobic and electrostatic interactions are important for the binding of group C N. meningitidis PS to the ELISA plate, while charge interactions seem to be sufficient for binding the more negatively charged H. influenzae type b PS.

L6 ANSWER 4 OF 5 SCISEARCH COPYRIGHT (c) 2006 The Thomson Corporation on STN

ACCESSION NUMBER: 1994:313652 SCISEARCH

THE GENUINE ARTICLE: NL733

TITLE: IDENTIFICATION OF LACTOFERRIN-BINDING PROTEINS FROM

TREPONEMA-PALLIDUM SUBSPECIES PALLIDUM AND

TREPONEMA-DENTICOLA

STAGGS T M (Reprint); GREER M K; BASEMAN J B; HOLT S AUTHOR:

C; TRYON V V

UNIV TEXAS, HLTH SCI CTR, DEPT MICROBIOL, SAN ANTONIO, CORPORATE SOURCE:

TX 78284; UNIV TEXAS, HLTH SCI CTR, DEPT PERIODONT,

SAN ANTONIO, TX 78284

COUNTRY OF AUTHOR: USA

SOURCE: MOLECULAR MICROBIOLOGY, (MAY 1994) Vol. 12, No. 4, pp.

613-619.

ISSN: 0950-382X.

BLACKWELL SCIENCE LTD, OSNEY MEAD, OXFORD, OXON, PUBLISHER:

ENGLAND OX2 OEL.

DOCUMENT TYPE: Article; Journal

FILE SEGMENT: LIFE LANGUAGE: English

REFERENCE COUNT: 33

ENTRY DATE: Entered STN: 1994

Last Updated on STN: 1994

\*ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS\*

Lactoferrin-binding or -associated proteins were identified in AB Treponema pallidum subspecies pallidum and Treponema denticola by affinity column chromatography using human lactoferrin and detergent-solubilized, radiolabelled spirochaetes. Two discrete polypeptides of T. pallidum with masses of 45 and 40 kDa and a broad band from 29-34 kDa exhibited association with human apo- and partially ferrated lactoferrin. T. denticola produced two proteins that associated with a lactoferrin affinity matrix (50 and 35 kDa). T. pallidum and T. denticola did not associate with soluble, human transferrin in parallel experiments. Soluble human lactoferrin competed with all lactoferrin-associated proteins from T. pallidum and T. denticola in competitive-binding assays. However, the T. denticola proteins dissociated from a lactoferrin-affinity matrix in the presence of differing concentrations of unlabelled, soluble lactoferrin competitor. Treatment with phospholipase p altered migration of the diffuse 29-34 kDa band of T. pallidum suggesting that the polypeptide was lipid-modified. Each of the lactoferrin-binding proteins from T. pallidum and T. denticola reacted with pooled rabbit syphilitic antisera. The lactoferrin-binding proteins of T. pallidum reacted with human sera from patients at all stages of syphilis. In addition, a monoclonal antibody generated against the  $45\ \mathrm{kDa}$  polypeptide of T. pallidum

crossreacted with the 29-34 kDa protein. ANSWER 5 OF 5 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on

L6

STN

ACCESSION NUMBER: 1986:242880 BIOSIS

DOCUMENT NUMBER: PREV198682007384; BA82:7384

INTERRELATIONSHIPS BETWEEN ALDEHYDE DEHYDROGENASE OF TITLE:

ACINETOBACTER-CALCOACETICUS AND MEMBRANE LIPIDS II. RECONSTITUTION IN ARTIFICIAL MEMBRANE VESICLES.

AURICH H [Reprint author]; BERGMANN R; LASCH J; KOELSCH AUTHOR(S):

R; SORGER H

INST BIOCHEMIE, BEREICH MED, MARTIN-LUTHER-UNIV, CORPORATE SOURCE:

HALLE-WITTENBERG, DDR-4020 HALLE, HOLLYSTR 1

Journal of Basic Microbiology, (1985) Vol. 25, No. 10, SOURCE:

pp. 631-636.

CODEN: JBMIEQ. ISSN: 0233-111X.

DOCUMENT TYPE: Article FILE SEGMENT: BA LANGUAGE: GERMAN

> Shears 571-272-2528

ENTRY DATE: Entered STN: 7 Jun 1986

Last Updated on STN: 7 Jun 1986

AB Purified aldehyde dehydrogenase (NADP+-dependent) of intracytoplasmic membranes of Acinetobacter calcoaceticus could be incorporated from micelles formed during the purification procedure into liposomal membranes. Both the cholate dilution method and the ultrasonication method were suitable to produce enzyme liposomes. In unilamellar liposomes produced by phosphatidyl choline, the enzyme activity decreased to 1% (or less) of the original activity. In contrast, about 10% of the original activity could be preserved in unilamellar liposomes prepared from bacterial phospholipids. The destruction of the enzyme liposomes induced by detergents (lauroyl sarcosinate) was followed by measuring the wavelength dependence of turbidity, which allowed us to draw conclusions on size and stability of the particles in the suspension. In addition these measurements demonstrated that decanal and NADP+ did not destroy the liposomal structure at concentrations necessary for the determination of enzyme activity. The liposomal enzyme was inactivated to a lesser degree by proteinase K than the micellar enzyme. Both phospholipase A2 and D inactivated the enzyme incorporated into the liposomal membranes to about 50%. After treatment with phospholipase A2, the enzyme could be reactivated by bacterial phospholipids. After treatment with phospholipase D, no reactivation was possible by bacterial phospholipids.

FILE 'CAPLUS' ENTERED AT 10:47:01 ON 03 MAY 2006

2 SEA ABB=ON PLU=ON PLD AND NEISSER? L7

O SEA ABB=ON PLU=ON L7 NOT L4 L8

> FILE 'MEDLINE, BIOSIS, EMBASE, WPIDS, CONFSCI, SCISEARCH, JICST-EPLUS, JAPIO' ENTERED AT 10:47:23 ON 03 MAY 2006

L9

9 SEA ABB=ON PLU=ON L7 4 SEA ABB=ON PLU=ON L9 NOT L5 L10

L114 DUP REM L10 (0 DUPLICATES REMOVED)

1 SEA ABB=ON PLU=ON L11 AND (POLYPEPTIDE OR PEPTIDE OR L12 PROTEIN OR POLYPROTEIN)

L13 O SEA ABB=ON PLU=ON L12 AND (VACCIN? OR IMMUNIS? OR IMMUNIZ?)

FILE 'MEDLINE' ENTERED AT 10:49:32 ON 03 MAY 2006

FILE LAST UPDATED: 2 MAY 2006 (20060502/UP). FILE COVERS 1950 TO DATE.

On December 11, 2005, the 2006 MeSH terms were loaded.

The MEDLINE reload for 2006 is now (26 Feb.) available. For details on the 2006 reload, enter HELP RLOAD at an arrow prompt (=>). See also:

http://www.nlm.nih.gov/mesh/

http://www.nlm.nih.gov/pubs/techbull/nd04/nd04 mesh.html

http://www.nlm.nih.gov/pubs/techbull/nd05/nd05\_med\_data\_changes.html

http://www.nlm.nih.gov/pubs/techbull/nd05/nd05\_2006\_MeSH.html

OLDMEDLINE is covered back to 1950.

MEDLINE thesauri in the /CN, /CT, and /MN fields incorporate the MeSH 2006 vocabulary.

> Shears 571-272-2528 :

This file contains CAS Registry Numbers for easy and accurate substance identification.

L14 O SEA FILE=MEDLINE ABB=ON PLU=ON (PHOSPHOLIPASE D AND

NEISSERIA)/CT

L15 6 SEA FILE=MEDLINE ABB=ON PLU=ON (PHOSPHOLIPASE D AND

BACTERIA)/CT

L15 ANSWER 1 OF 6 MEDLINE on STN

ACCESSION NUMBER: 2005430216 MEDLINE DOCUMENT NUMBER: PubMed ID: 16096028

TITLE: Non-HKD phospholipase D enzymes: new players in

phosphatidic acid signaling?.

AUTHOR: Zambonelli Carlo; Roberts Mary F

CORPORATE SOURCE: Merkert Chemistry Center, Boston College, Chestnut

Hill, Massachusetts 02467, USA.

SOURCE: Progress in nucleic acid research and molecular

biology, (2005) Vol. 79, pp. 133-81. Ref: 206

Journal code: 0102753. ISSN: 0079-6603.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

General Review; (REVIEW)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200508

ENTRY DATE: Entered STN: 15 Aug 2005

Last Updated on STN: 31 Aug 2005 Entered Medline: 30 Aug 2005

ED Entered STN: 15 Aug 2005

Last Updated on STN: 31 Aug 2005 Entered Medline: 30 Aug 2005

L15 ANSWER 2 OF 6 MEDLINE on STN

ACCESSION NUMBER: 2004324113 MEDLINE DOCUMENT NUMBER: PubMed ID: 15225639

TITLE: A distant evolutionary relationship between GPI-specific phospholipase D and bacterial

phosphatidylcholine-preferring phospholipase C.

AUTHOR: Rigden Daniel J

CORPORATE SOURCE: School of Biological Sciences, University of Liverpool,

Crown Street, Liverpool L69 7ZB, UK.. drigden@liv.ac.uk

SOURCE: FEBS letters, (2004 Jul 2) Vol. 569, No. 1-3, pp.

229-34.

Journal code: 0155157. ISSN: 0014-5793.

PUB. COUNTRY: Netherlands

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200408

ENTRY DATE: Entered STN: 1 Jul 2004

Last Updated on STN: 26 Aug 2004 Entered Medline: 25 Aug 2004

ED Entered STN: 1 Jul 2004

Last Updated on STN: 26 Aug 2004 Entered Medline: 25 Aug 2004

AB In eukaryotes some surface proteins are attached to the plasma membrane by a glycosylphosphatidylinositol (GPI) anchor. A

GPI-specific phospholipase D (GPI-PLD) activity has been characterized

and implicated in the regulation of anchoring, thereby influencing the dispersal of anchored proteins or their maintenance on the cell surface, and possibly in cell signalling. Despite its biological and medical importance, little is known of the structure of GPI-PLD. Here, a distant relationship between the catalytic domains of GPI-PLD and some bacterial phospholipases C is demonstrated. A model of the GPI-PLD catalytic site sheds light on catalysis and highlights possibilities for design of improved and more specific GPI-PLD inhibitors. The databases contain hitherto unnoticed close homologues of GPI-PLD from yeast and Dictyostelium discoideum.

L15 ANSWER 3 OF 6 MEDLINE on STN
ACCESSION NUMBER: 2001206925 MEDLINE
DOCUMENT NUMBER: PubMed ID: 11239820

TITLE: Cloning and direct G-protein regulation of

phospholipase D from tobacco.

AUTHOR: Lein W; Saalbach G

CORPORATE SOURCE: Institute of Plant Genetics and Crop Plant Research,

Corrensstrasse 3, D-06466, Gatersleben, Germany.

SOURCE: Biochimica et biophysica acta, (2001 Feb 26) Vol. 1530,

No. 2-3, pp. 172-83.

Journal code: 0217513. ISSN: 0006-3002.

PUB. COUNTRY: Netherlands

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200104

ENTRY DATE: Entered STN: 17 Apr 2001

Last Updated on STN: 17 Apr 2001 Entered Medline: 12 Apr 2001

ED Entered STN: 17 Apr 2001

Last Updated on STN: 17 Apr 2001 Entered Medline: 12 Apr 2001

AB Phospholipase D (PLD) and heterotrimeric G-proteins are involved in plant signal transduction pathways at the plasma membrane. There is evidence suggesting that PLD acts downstream from G-proteins, but a direct interaction of specific members has not been shown. In the present paper, a PLD cDNA clone was isolated from tobacco, expressed as a GST fusion in bacteria, and the recombinant protein was purified by glutathione affinity. Its enzymatic properties identified it as an alpha-type PLD. The alpha-subunit of a G-protein from tobacco was isolated in a similar way. Both proteins were functional in biochemical assays. When the G-protein was included in the PLD assay, a strong dosage-dependent inhibition of the PLD activity was observed. Different control proteins did not exhibit this inhibitory effect. When GST-NtGPalphal was activated by incubation with GTPgammaS the inhibitory activity was greatly reduced. These results provide a first indication for a direct regulation of PLDalpha by a

L15 ANSWER 4 OF 6 MEDLINE on STN ACCESSION NUMBER: 96303814 MEDLINE DOCUMENT NUMBER: PubMed ID: 8732763

TITLE: A novel family of phospholipase D homologues that

heterotrimeric G-protein alpha-subunit in plants.

includes phospholipid synthases and putative

endonucleases: identification of duplicated repeats and

potential active site residues.

AUTHOR: Ponting C P; Kerr I D

CORPORATE SOURCE: Fibrinolysis Research Unit, University of Oxford,

United Kingdom.. chris@biop.ox.ac.uk

SOURCE: Protein science: a publication of the Protein Society,

(1996 May) Vol. 5, No. 5, pp. 914-22. Journal code: 9211750. ISSN: 0961-8368.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

OTHER SOURCE: GENBANK-R34925; GENBANK-R83570; GENBANK-T76232;

GENBANK-T88610; GENBANK-Z45777

ENTRY MONTH: 199702

ENTRY DATE: Entered STN: 6 Mar 1997

Last Updated on STN: 6 Mar 1997 Entered Medline: 21 Feb 1997

ED Entered STN: 6 Mar 1997

Last Updated on STN: 6 Mar 1997 Entered Medline: 21 Feb 1997

AB Phosphatidylcholine-specific phospholipase D (PLD) enzymes catalyze hydrolysis of phospholipid phosphodiester bonds, and also transphosphatidylation of phospholipids to acceptor alcohols. Bacterial and plant PLD enzymes have not been shown previously to be homologues or to be homologous to any other protein. Here we show, using sequence analysis methods, that bacterial and plant PLDs show significant sequence similarities both to each other, and to two other classes of phospholipid-specific enzymes, bacterial cardiolipin synthases, and eukaryotic and bacterial phosphatidylserine synthases, indicating that these enzymes form an homologous family. This family is suggested also to include two Poxviridae proteins of unknown function (p37K and protein K4), a bacterial endonuclease (nuc), an Escherichia coli putative protein (0338) containing an N-terminal domain showing similarities with helicase motifs V and VI, and a Synechocystis sp. putative protein with a C-terminal domain likely to possess a DNA-binding function. Surprisingly, four regions of sequence similarity that occur once in nuc and o338, appear twice in all other homologues, indicating that the latter molecules are bi-lobed, having evolved from an ancestor or ancestors that underwent a gene duplication and fusion event. It is suggested that, for each of these enzymes, conserved histidine, lysine, aspartic acid, and/or asparagine residues may be involved in a two-step ping pong mechanism involving an enzyme-substrate intermediate.

L15 ANSWER 5 OF 6 MEDLINE on STN ACCESSION NUMBER: 96102003 MEDLINE DOCUMENT NUMBER: PubMed ID: 8530346

TITLE: Human ADP-ribosylation factor-activated

phosphatidylcholine-specific phospholipase D defines a

new and highly conserved gene family.

AUTHOR: Hammond S M; Altshuller Y M; Sung T C; Rudge S A; Rose

K; Engebrecht J; Morris A J; Frohman M A

CORPORATE SOURCE: Department of Pharmacological Sciences, State

University of New York, Stony Brook 11794-8651, USA.

CONTRACT NUMBER: GM4863903 (NIGMS)

GM50388 (NIGMS) HD29758 (NICHD)

+ -

SOURCE: The Journal of biological chemistry, (1995 Dec 15) Vol.

270, No. 50, pp. 29640-3.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

OTHER SOURCE: GENBANK-D27058; GENBANK-D33536; GENBANK-G00778; GENBANK-L33686; GENBANK-T76232; GENBANK-T88610;

GENBANK-U38545; GENBANK-X28256; GENBANK-Z18424;

GENBANK-Z33674

ENTRY MONTH: 199601

ENTRY DATE: Entered STN: 20 Feb 1996

Last Updated on STN: 3 Mar 2000 Entered Medline: 26 Jan 1996

ED Entered STN: 20 Feb 1996

Last Updated on STN: 3 Mar 2000 Entered Medline: 26 Jan 1996

AB Activation of phosphatidylcholine-specific phospholipase D (PLD) has been implicated as a critical step in numerous cellular pathways, including signal transduction, membrane trafficking, and the regulation of mitosis. We report here the identification of the first human PLD cDNA, which defines a new and highly conserved gene family. Characterization of recombinant human PLD1 reveals that it is membrane-associated, selective for phosphatidylcholine, stimulated by phosphatidylinositol 4,5-bisphosphate, activated by the monomeric G-protein ADP-ribosylation factor-1, and inhibited by oleate. PLD1 likely encodes the gene product responsible for the most widely studied endogenous PLD activity.

L15 ANSWER 6 OF 6 MEDLINE on STN
ACCESSION NUMBER: 82087704 MEDLINE
DOCUMENT NUMBER: PubMed ID: 6274146

TITLE: Enzymatic hydrolysis by bacterial phospolipases C and D

of immobilized radioactive sphingomyelin and

phosphatidylcholine.
Malmqvist T; Mollby R

SOURCE: Acta pathologica et microbiologica Scandinavica.

Section B, Microbiology, (1981 Oct) Vol. 89, No. 5, pp.

363-7.

Journal code: 7508472. ISSN: 0105-0656.

PUB. COUNTRY: Denmark

AUTHOR:

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 198202

ENTRY DATE: Entered STN: 16 Mar 1990

Läst Updated on STN: 16 Mar 1990 Entered Medline: 12 Feb 1982

ED Entered STN: 16 Mar 1990

Last Updated on STN: 16 Mar 1990 Entered Medline: 12 Feb 1982

AB An assay system for phospholipases C has been described with sphingomyelin immobilized to octyl-Sepharose CL-4B as substrate. The immobilization procedure was further developed and used with [14 C-choline]-sphingomyelin and [14C-choline] phosphatidylcholine (lecithin). These immobilized radioactive phospholipids made the enzymatic assays easier to perform and made it possible to increase the sensitivity. Furthermore, since release of the choline part instead of the phosphate part of the substrate molecule was measured, it was possible to use this assay for phospholipase D as well. The enzyme characteristics of phospholipase D from Corynebacterium ovis were compared in this test system with those of three phospholipases C

(from Clostridium perfringens, Bacillus cereus and Staphylococcus aureus) with respect to hydrolysing capacities and optimal ion concentrations.

FILE 'USPATFULL' ENTERED AT 10:50:39 ON 03 MAY 2006 CA INDEXING COPYRIGHT (C) 2006 AMERICAN CHEMICAL SOCIETY (ACS)

FILE COVERS 1971 TO PATENT PUBLICATION DATE: 2 May 2006 (20060502/PD)

FILE LAST UPDATED: 2 May 2006 (20060502/ED) HIGHEST GRANTED PATENT NUMBER: US7039955

HIGHEST APPLICATION PUBLICATION NUMBER: US2006090232 CA INDEXING IS CURRENT THROUGH 2 May 2006 (20060502/UPCA)

ISSUE CLASS FIELDS (/INCL) CURRENT THROUGH: 2 May 2006 (20060502/PD)

REVISED CLASS FIELDS (/NCL) LAST RELOADED: Feb 2006

USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Feb 2006

L1	154 SEA FILE=REGISTRY ABB=ON PLU=ON PHOSPHOLIPASE D ?/CN
L2	4852 SEA FILE=CAPLUS ABB=ON PLU=ON L1 OR (PHOSPHOLIPASE OR
	PHOSPHO LIPASE OR LECITHINASE) (1W) D OR (PHOSPHATIDYLCHOLIN
	OR PHOSPHATIDYL CHOLINE) (W) (PHOSPHOHYDROLASE OR PHOSPHO
	HYDROLASE)
Ļ19	564 SEA FILE-USPATFULL ABB=ON PLU=ON (L2 OR PLD)(S)(POLYPEPT
	DE OR PEPTIDE OR PROTEIN OR POLYPROTEIN)
L20	23 SEA FILE=USPATFULL ABB=ON PLU=ON L19(L)NEISSER?
L21	22 SEA FILE=USPATFULL ABB=ON PLU=ON L20(L)(VACCIN? OR
	IMMUNIS? OR IMMUNIZ?)

L21 ANSWER 1 OF 22 USPATFULL on STN

ACCESSION NUMBER: 2006:80413 USPATFULL

TITLE: Single-stranded nucleic acid template-mediated recombination and nucleic acid fragment isolation

- Affholter, Joseph A., Zephyr Cove, NV, UNITED INVENTOR(S):

STATES

Cox, Anthony, Mountain View, CA, UNITED STATES Ness, Jon E., Redwood City, CA, UNITED STATES

Carr, Brian, Raleigh, NC, UNITED STATES

PATENT ASSIGNEE(S): Maxygen, Inc. (U.S. corporation)

DATE NUMBER KIND US 2006068406 A1 20060330 US 2005-47380 A1 20050131 (11) PATENT INFORMATION: APPLICATION INFO.: Continuation of Ser. No. US 2000-721507, filed on RELATED APPLN. INFO.: 22 Nov 2000, ABANDONED Continuation of Ser. No. US 2000-656549, filed on 6 Sep 2000, ABANDONED

			NUMBER	DATE	
PRIORITY	INFORMATION:	US	2000-185244P	20000228	(60)
		US	2000-185815P	20000229	(60)
		US	2000-186247P	20000301	(60)
		US	2000-186482P	20000302	(60)
DOCUMENT	myne.	TT4-3	. 7		

DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: MAXYGEN, INC., INTELLECTUAL PROPERTY DEPARTMENT,

515 GALVESTON DRIVE, RED WOOD CITY, CA, 94063, US

NUMBER OF CLAIMS: 22 EXEMPLARY CLAIM: 1-43

NUMBER OF DRAWINGS: 8 Drawing Page(s)

6266 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Methods mediated by single-stranded nucleic acid templates, including utilizing single-stranded nucleic acid templates to isolate nucleic acid fragments and to recombine nucleic acid fragments. Methods include polymerase and polymerase-free recombination of nucleic acid fragments to generate chimeric nucleic acid sequences. Integrated systems and kits are also provided.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 2 OF 22 USPATFULL on STN

ACCESSION NUMBER: 2006:41437 USPATFULL

TITLE: 33 human secreted proteins

Soppet, Daniel R., Centreville, VA, UNITED STATES INVENTOR(S):

Moore, Paul A., North Bethesda, MD, UNITED STATES Shi, Yanggu, Gaithersburg, MD, UNITED STATES Ruben, Steven M., Brookeville, MD, UNITED STATES Rosen, Craig A., Laytonsville, MD, UNITED STATES LaFleur, David W., Washington, DC, UNITED STATES Olsen, Henrik S., Gaithersburg, MD, UNITED STATES Ebner, Reinhard, Gaithersburg, MD, UNITED STATES Florence, Kimberly, Rockville, MD, UNITED STATES Young, Paul, Gaithersburg, MD, UNITED STATES Komatsoulis, George, Silver Spring, MD, UNITED

STATES

Ni, Jian, Germantown, MD, UNITED STATES

PATENT ASSIGNEE(S): Human Genome Sciences, Inc., Rockville, MD, UNITED

STATES (U.S. corporation)

NUMBER KIND DATE

US 2006036089 PATENT INFORMATION: A1 20060216 US 2005-240769 A1 20051003 (11) APPLICATION INFO.:

Continuation of Ser. No. US 2001-997131, filed on RELATED APPLN. INFO.: 30 Nov 2001, PENDING Continuation of Ser. No. US 2000-628508, filed on 28 Jul 2000, ABANDONED

Continuation-in-part of Ser. No. WO 2000-US3062,

filed on 8 Feb 2000, PENDING

NUMBER DATE

US 1999-119468P 19990210 (60) PRIORITY INFORMATION:

DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT:

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, INTELLECTUAL PROPERTY

DEPT., 14200 SHADY GROVE ROAD, ROCKVILLE, MD,

20850, US

20 NUMBER OF CLAIMS: EXEMPLARY CLAIM: LINE COUNT: 17123

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to 33 novel human secreted proteins and isolated nucleic acids containing the coding regions of the genes encoding such proteins. Also provided are vectors, host cells, antibodies, and recombinant methods for producing human secreted proteins. The invention further relates to diagnostic and therapeutic methods useful for diagnosing and treating diseases, disorders, and/or conditions related to these novel human secreted

proteins.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 3 OF 22 USPATFULL on STN

ACCESSION NUMBER: 2006:40224 USPATFULL

TITLE: Immunogenic compositions for Chlamydia trachomatis

INVENTOR(S): Grandi, Guido, Milano, ITALY
Ratti, Guilio, Siena, ITALY

Bonci, Alessandra, Siena, ITALY

Finco, Oretta, Castelnuovo Berardenga, ITALY

PATENT ASSIGNEE(S): Chiron Corporation, Emeryville, CA, UNITED STATES

(U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: US 2006034871 A1 20060216 APPLICATION INFO.: US 2004-18868 A1 20041222 (11)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. WO 2004-US20491,

filed on 25 Jun 2004, PENDING

DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: Chiron Corporation, Intellectual Property - R440,

P.O. Box 8097, Emeryville, CA, 94662-8097, US

NUMBER OF CLAIMS: 45 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 3 Drawing Page(s)

LINE COUNT: 9932

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The invention relates to immunogenic compositions comprising combinations of Chlamydia trachomatis antigens and their use in vaccines. The composition may comprise at least two components, one component of which comprises Chlamydia trachomatis antigens for eliciting a Chlamydia trachomatis specific TH1 immune response and another component of which comprises antigens for eliciting a Chlamydia trachomatis specific TH2 immune response. The invention further relates to an immunogenic composition comprising a Chlamydia trachomatis Type III secretion system (TTSS) regulatory protein and a Chlamydia trachomatis Type III secretion system (TTSS) secreted protein or a fragment thereof. The invention further relates to the use of combinations of adjuvants for use with antigens associated with a sexually transmissible disease, such as Chlamydia trachomatis antigens. Preferred adjuvant combinations include mineral salts, such as aluminium salts and oligonucleotides comprising a CpG motif. The invention further provides a combination of Chlamydia trachomatis antigens comprising a Chlamydia trachomatis antigen that is conserved over at least two serovars.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 4 OF 22 USPATFULL on STN

ACCESSION NUMBER: 2005:234340 USPATFULL

TITLE: Alloiococcus otitidis open reading frames (orfs)

encoding polypeptide antigens, immunogenic

compositions and uses thereof

McMichael, John Calhoun, Rochester, NY, UNITED INVENTOR(S):

Zagursky, Robert John, Victor, NY, UNITED STATES Fletcher, Leah Diane, Geneseo, NY, UNITED STATES

DATE NUMBER KIND \_\_\_\_\_\_\_ US 2005203280 A1 20050915 US 2003-501282 A1 20021125 PATENT INFORMATION: (10) APPLICATION INFO.: 20021125 WO 2002-US36123 20021125 20040709 PCT 371 date

> NUMBER DATE

US 2001-60333777 20011129 PRIORITY INFORMATION:

US 2003-60426742 20021118

DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT:

LEGAL REPRESENTATIVE: WYETH, PATENT LAW GROUP, 5 GIRALDA FARMS, MADISON,

NJ, 07940, US

NUMBER OF CLAIMS: 107 EXEMPLARY CLAIM: LINE COUNT: 36418

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to the complete genomic sequence of AB Gram-positive bacterium, Alloiococcus otitidis. The present invention also relates to polynucleotide sequences encoding polypeptides of Alloiococcus otitidis. In particular, the invention relates to antigenic polypeptides encoded by the Alloiococcus otitidis open reading frames (ORFs), and to their use in immunogenic compositions, therapeutics, diagnostics and the like.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 5 OF 22 USPATFULL on STN

2005:151277 USPATFULL ACCESSION NUMBER:

TITLE: Compositions and methods for treating and

diagnosing irritable bowel syndrome

Pasricha, Pankaj, Houston, TX, UNITED STATES INVENTOR(S): Shenoy, Mohan, Galveston, TX, UNITED STATES

Winston, John, League City, TX, UNITED STATES

NUMBER KIND DATE \_\_\_\_\_\_ US 2005130189 A1 20050616 US 2004-923035 A1 20040823 (10) PATENT INFORMATION: APPLICATION INFO.:

NUMBER DATE \_\_\_\_\_

PRIORITY INFORMATION: US 2003-496716P 20030821 (60)

Utility DOCUMENT TYPE: APPLICATION FILE SEGMENT:

LEGAL REPRESENTATIVE: Supervisor, Patent Prosecuting Services, PIPER

RUDNICK LLP, 1200 Nineteenth Street, N.W.,

Washington, DC, 20036-2412, US

NUMBER OF CLAIMS: 31

EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 3 Drawing Page(s)

LINE COUNT:

9702

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB

Compositions and methods for diagnosing and treating CVH and CVH-associated disorders are disclosed. Genes differentially expressed in CVH tissues relative to normal tissues are identified. The genes and the gene products (i.e., the polynucleotides transcribed from and polypeptides encoded by the genes) can be used as markers of CVH. The genes and the gene products can also be used to screen agents that modulate the gene expression or the activities of the gene products.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 6 OF 22 USPATFULL on STN

ACCESSION NUMBER: 2005:75161 USPATFULL

TITLE:

143 human secreted proteins

INVENTOR(S):

Rosen, Craig A., Laytonsville, MD, UNITED STATES Ruben, Steven M., Brookeville, MD, UNITED STATES Moore, Paul A., North Bethesda, MD, UNITED STATES Young, Paul E., Gaithersburg, MD, UNITED STATES Komatsoulis, George, Silver Spring, MD, UNITED

Birse, Charles E., North Potomac, MD, UNITED STATES Duan, Roxanne D., Gaithersburg, MD, UNITED STATES Florence, Kimberly A., Rockville, MD, UNITED STATES Soppet, Daniel R., Centreville, VA, UNITED STATES

PATENT ASSIGNEE(S):

Human Genome Sciences, Inc., Rockville, MD (U.S.

corporation)

NUMBER KIND DATE \_\_\_\_\_\_ US 2005064458 A1 20050324 US 2004-863332 A1 20040609 (10)

PATENT INFORMATION: APPLICATION INFO.:

RELATED APPLN. INFO.: -

Continuation of Ser. No. US 2001-986480, filed on 8 Nov 2001, ABANDONED Continuation-in-part of Ser.

No. WO 2000-US12788, filed on 11 May 2000, PENDING

NUMBER DATE \_\_\_\_\_

PRIORITY INFORMATION:

US 1999-134068P 19990513 (60)

DOCUMENT TYPE:

Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, INTELLECTUAL PROPERTY

DEPT., 14200 SHADY GROVE ROAD, ROCKVILLE, MD, 20850

NUMBER OF CLAIMS:

EXEMPLARY CLAIM:

LINE COUNT:

26589

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to novel human secreted proteins and isolated nucleic acids containing the coding regions of the genes encoding such proteins. Also provided are vectors, host cells, antibodies, and recombinant methods for producing human secreted proteins. The invention further relates to diagnostic and therapeutic methods useful for diagnosing and treating diseases, disorders, and/or conditions related to these novel human secreted proteins.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 7 OF 22 USPATFULL on STN

ACCESSION NUMBER: 2004:320569 USPATFULL

Vaccine and compositions for the prevention and TITLE:

treatment of neisserial infections

Apicella, Michael A., Solon, IA, UNITED STATES INVENTOR(S):

Edwards, Jennifer L., Iowa City, IA, UNITED STATES

KIND DATE NUMBER \_\_\_\_\_\_

PATENT INFORMATION: US 2004253222 A1 20041216 US 2003-665990 A1 20030919 (10)

APPLICATION INFO.:

Continuation-in-part of Ser. No. US 2003-621184, RELATED APPLN. INFO.: filed on 15 Jul 2003, PENDING Continuation-in-part of Ser. No. US 2002-66551, filed on 31 Jan 2002,

PENDING

NUMBER DATE \_\_\_\_\_

PRIORITY INFORMATION:

US 2001-266070P 20010131 (60) US 2001-310356P 20010806 (60) US 2001-344452P 20011023 (60)

DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION
LEGAL REPRESENTATIVE: FISH & RICHARDSON P.C., 3300 DAIN RAUSCHER PLAZA,

60 SOUTH SIXTH STREET, MINNEAPOLIS, MN, 55402

60 24 NUMBER OF CLAIMS: 1 EXEMPLARY CLAIM:

EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 23 Drawing Page(s)

LINE COUNT: 6288

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention is directed to novel polypeptides,

polynucleotides and vaccines for use against Neisseria gonorrhoeae colonization or infection and/or Neisseria meningitidis colonization

or infection. The vaccines contain an immunogenic amount of a

neisserial protein.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 8 OF 22 USPATFULL on STN

ACCESSION NUMBER: 2004:141216 USPATFULL

Nucleic acid sequences relating to Candida albicans TITLE:

for diagnostics and therapeutics

Weinstock, Keith G., Westborough, MA, United States INVENTOR(S):

Bush, David, Somerville, MA, United States

PATENT ASSIGNEE(S): Genome Therapeutics Corporation, Waltham, MA,

United States (U.S. corporation)

NUMBER KIND \_\_\_\_\_

US 6747137 B1 20040608 US 1999-248796 19990212 PATENT INFORMATION: 19990212 (9) APPLICATION INFO.:

NUMBER DATE

\_\_\_\_\_

US 1998-96409P 19980813 (60) US 1998-74725P 19980213 (60) PRIORITY INFORMATION:

DOCUMENT TYPE: Utility

FILE SEGMENT:

GRANTED

PRIMARY EXAMINER:

Marschel, Ardin H.

LEGAL REPRESENTATIVE: Genome Therapeutics Corporation

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

12

NUMBER OF DRAWINGS:

0 Drawing Figure(s); 0 Drawing Page(s)

LINE COUNT: 36816

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The invention provides isolated polypeptide and nucleic acid sequences derived from Candida albicans that are useful in diagnosis and therapy of pathological conditions; antibodies against the polypeptides; and methods for the production of the polypeptides. The invention also provides methods for the detection, prevention and treatment of pathological conditions resulting from fungal infection.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 9 OF 22 USPATFULL on STN

ACCESSION NUMBER:

2004:63731 USPATFULL

TITLE:

INVENTOR(S):

Novel nucleic acids and secreted polypeptides Tang, Y. Tom, San Jose, CA, UNITED STATES Yang, Yonghong, San Jose, CA, UNITED STATES Weng, Gezhi, Piedmont, CA, UNITED STATES Zhang, Jie, Campbell, CA, UNITED STATES Ren, Feiyan, Cupertino, CA, UNITED STATES Xue, Aidong, Sunnyvale, CA, UNITED STATES Wang, Jian-Rui, Cupertino, CA, UNITED STATES Wehrman, Tom, Stanford, CA, UNITED STATES Ghosh, Malabika J., Sunnyvale, CA, UNITED STATES Wang, Dunrui, Poway, CA, UNITED STATES

Zhao, Qing A., San Jose, CA, UNITED STATES Wang, Zhiwei, Sunnyvale, CA, UNITED STATES

NUMBER	KIND	DATE

RELATED APPLN. INFO.:

PATENT INFORMATION: US 2004048249 A1 20040311 APPLICATION INFO.: US 2002-112944 A1 20020328 20020328 (10)

Continuation-in-part of Ser. No. US 2000-488725, filed on 21 Jan 2000, PENDING Continuation-in-part of Ser. No. US 2000-491404, filed on 25 Jan 2000, ABANDONED Continuation-in-part of Ser. No. US 2000-496914, filed on 3 Feb 2000, ABANDONED Continuation-in-part of Ser. No. US 2000-515126,

filed on 28 Feb 2000, ABANDONED Continuation-in-part of Ser. No. US 2000-519705, filed on 7 Mar 2000, ABANDONED Continuation-in-part of Ser. No. US 2000-540217, filed on 31 Mar 2000, ABANDONED Continuation-in-part of Ser. No. US 2000-552929, filed on 18 Apr 2000, ABANDONED Continuation-in-part of Ser. No. US 2000-577408,

filed on 18 May 2000, ABANDONED

NUMBER DATE \_\_\_\_\_\_

PRIORITY INFORMATION:

US 2001-306971P 20010721 (60)

DOCUMENT TYPE: FILE SEGMENT:

Utility APPLICATION

LEGAL REPRESENTATIVE:

Luisa Biogornia, HYSEQ, INC., 670 Almanor Avenue,

Sunnyvale, CA, 94085

NUMBER OF CLAIMS: 26 EXEMPLARY CLAIM: LINE COUNT: 23809

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention provides novel nucleic acids, novel polypeptide sequences encoded by these nucleic acids and uses

thereof.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 10 OF 22 USPATFULL on STN

2004:20717 USPATFULL ACCESSION NUMBER:

TITLE: Rice promoters for regulation of plant expression

Budworth, Paul, San Diego, CA, UNITED STATES INVENTOR(S): Moughamer, Todd, San Diego, CA, UNITED STATES

Briggs, Steven P., Del Mar, CA, UNITED STATES Cooper, Bret, La Jolla, CA, UNITED STATES Glazebrook, Jane, San Diego, CA, UNITED STATES Goff, Stephen Arthur, Encinitas, CA, UNITED STATES

Katagiri, Fumiaki, San Diego, CA, UNITED STATES

Kreps, Joel, Carlsbad, CA, UNITED STATES

Provart, Nicholas, Toronto, CANADA

Ricke, Darrell, San Diego, CA, UNITED STATES

Zhu, Tong, San Diego, CA, UNITED STATES

NUMBER KIND DATE US 2004016025 A1 20040122 US 2002-260238 A1 20020926 PATENT INFORMATION: APPLICATION INFO.:

20020926 (10)

NUMBER DATE \_\_\_\_\_

US 2001-325448P 20010926 (60) US 2001-325277P 20010926 (60) US 2002-370620P 20020404 (60) PRIORITY INFORMATION:

DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT:

LEGAL REPRESENTATIVE: James E. Butler, Torrey Mesa Research Institute,

3115 Merryfield Row, San Diego, CA, 92121

NUMBER OF CLAIMS: 77 EXEMPLARY CLAIM: 1 18818 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The invention provides a method to identify a plurality of plant promoters having a particular characteristic as well as the sequence

of promoters having one of those characteristics.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 11 OF 22 USPATFULL on STN

ACCESSION NUMBER: 2004:12649 USPATFULL TITLE: Anti-pathogen treatments

INVENTOR(S): Rider, Todd H., Littleton, MA, UNITED STATES PATENT ASSIGNEE(S): Massachusetts Institute of Technology, Cambridge,

MA (U.S. corporation)

NUMBER KIND DATE 

A1 PATENT INFORMATION: US 2004009167 20040115

US 2003-361208 APPLICATION INFO.: A1 20030207 (10)

> DATE NUMBER -----

PRIORITY INFORMATION:

US 2002-355359P 20020207 (60) US 2002-355022P 20020207 (60) US 2002-432386P 20021210 (60)

Utility DOCUMENT TYPE: FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: HAMILTON, BROOK, SMITH & REYNOLDS, P.C., 530

VIRGINIA ROAD, P.O. BOX 9133, CONCORD, MA,

01742-9133

NUMBER OF CLAIMS: 26 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 86 Drawing Page(s)

LINE COUNT: 9654

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Chimeric molecules that contain at least one pathogen-detection domain and at least one effector domain, and their methods of use in preventing or treating a pathogen infection in a cell or organism are described. The pathogen-detection domain and effector domain of the chimeric molecules are domains not typically found in nature to be associated together. Agents are also described herein having at least one pathogen-interacting molecular structure and at least one effector-mediating molecular structure, the agent being one that is non-naturally-occurring in a cell. The methods of prevention and treatment described herein are effective for a broad spectrum of pathogens and exhibit little or no toxic side-effects. Assays for the detection of a pathogen, pathogen component, or product produced or induced by a pathogen, are also provided.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 12 OF 22 USPATFULL on STN

2003:334942 USPATFULL ACCESSION NUMBER:

TITLE: Immunogenic peptides, and method of identifying

INVENTOR(S): Katritch, Vsevolod, San Diego, CA, UNITED STATES

Bordner, Andrew, San Diego, CA, UNITED STATES Deans, Robert, Claremont, CA, UNITED STATES Sumner, Mary, San Diego, CA, UNITED STATES

NUMBER KIND DATE \_\_\_\_\_\_ PATENT INFORMATION:

US 2003235818 A1 20031225 US 2003-410647 A1 20030408 A1 20030408 (10) APPLICATION INFO.:

> NUMBER DATE \_\_\_\_\_

US 2002-371250P 20020408 (60) US 2002-371256P 20020408 (60) PRIORITY INFORMATION:

US 2002-373668P 20020417 (60)

DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION

LISA A. HAILE, J.D., PH.D., GRAY CARY WARE & LEGAL REPRESENTATIVE:

FREIDENRICH LLP, Suite 1100, 4365 Executive Drive,

San Diego, CA, 92121-2133

NUMBER OF CLAIMS: 118

EXEMPLARY CLAIM:

1 NUMBER OF DRAWINGS: 4 Drawing Page(s)

LINE COUNT:

\_ 3957

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Immunogenic peptides, polynucleotides encoding immunogenic peptides, antibodies that selectively bind immunogenic peptides and methods of identifying immunogenic peptides are provided. The immunogenic peptides are representative of a structural element of a target protein. The methods of the invention are useful for identifying immunogenic peptides of a target protein having a known three dimensional structure, or of a target protein having a known amino acid sequence but an unknown three dimensional structure.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 13 OF 22 USPATFULL on STN

ACCESSION NUMBER:

2003:318635 USPATFULL

TITLE:

INVENTOR(S):

Novel nucleic acids and polypeptides Tang, Y. Tom, San Jose, CA, UNITED STATES Yang, Yonghong, San Jose, CA, UNITED STATES Wang, Zhiwei, Sunnyvale, CA, UNITED STATES

Weng, Gezhi, Piedmont, CA, UNITED STATES Ma, Yunqing, Santa Clara, CA, UNITED STATES DATE NUMBER KIND

-----US 2003224379 A1 20031204 US 2002-243552 A1 20020912 (10) PATENT INFORMATION:

APPLICATION INFO.:

RELATED APPLN. INFO.:

Continuation-in-part of Ser. No. WO 2000-US35017, filed on 22 Dec 2000, PENDING Continuation-in-part of Ser. No. US 2000-552317, filed on 25 Apr 2000, ABANDONED Continuation-in-part of Ser. No. US 2000-488725, filed on 21 Jan 2000, PENDING

			NUMBER	DATE	
PRIORITY	INFORMATION:	WO	2001-US2623	20010125	
		WO	2001-US3800	20010205	
		WO	2001-US4927	20010226	
		WO	2001-US4941	20010305	
		WO	2001-US8631	20010330	
		WO	2001-US8656	20010416	
		WO	2001-US14827	20010516	
		US	2001-322511P	20010913	(60)

DOCUMENT TYPE: FILE SEGMENT:

Utility

APPLICATION LEGAL REPRESENTATIVE: Elena Quertermous, 675 Almanor Avenue, Sunnyvale,

CA, 94085

NUMBER OF CLAIMS: 26 EXEMPLARY CLAIM: 1 LINE COUNT: 13810

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention provides novel nucleic acids, novel polypeptide sequences encoded by these nucleic acids and uses thereof.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 14 OF 22 USPATFULL on STN

ACCESSION NUMBER: 2003:240330 USPATFULL

TITLE: Nucleic acid and amino acid sequences relating to

Enterococcus faecalis for diagnostics and

therapeutics

Doucette-Stamm, Lynn A., 14 Flanagan Dr., INVENTOR(S):

Framingham, MA, United States 01701

Bush, David, 205 Holland St., Somerville, MA,

United States 02144

NUMBER KIND DATE \_\_\_\_\_\_ US 6617156 B1 20030909 US 1998-134000 19980813 PATENT INFORMATION:

19980813 (9) APPLICATION INFO.:

> NUMBER DATE \_\_\_\_\_\_

PRIORITY INFORMATION: US 1997-55778P 19970815 (60)

DOCUMENT TYPE: Utility FILE SEGMENT: GRANTED

PRIMARY EXAMINER: Mosher, Mary E.

LEGAL REPRESENTATIVE: Genome Therapeutics Corporation

NUMBER OF CLAIMS: 19 EXEMPLARY CLAIM: 1,5,14

NUMBER OF DRAWINGS: 0 Drawing Figure(s); 0 Drawing Page(s)

LINE COUNT: 13738

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The invention provides isolated polypeptide and nucleic acid sequences derived from Enterococcus faecalis that are useful in diagnosis and therapy of pathological conditions; antibodies against the polypeptides; and methods for the production of the polypeptides. The invention also provides methods for the detection,

prevention and treatment of pathological conditions resulting from

bacterial infection.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 15 OF 22 USPATFULL on STN

ACCESSION NUMBER: 2003:219631 USPATFULL

TITLE: Full-length human cDNAs encoding potentially

secreted proteins

Dumas Milne Edwards, Jean-Baptiste, Paris, FRANCE INVENTOR(S):

Bougueleret, Lydie, Petit Lancy, SWITZERLAND

Jobert, Severin, Paris, FRANCE

NUMBER KIND DATE \_\_\_\_\_\_ US 2003152921 A1 20030814 US 2001-876997 A1 20010608 PATENT INFORMATION:
APPLICATION INFO.: 20010608 (9)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 2000-731872,

filed on 7 Dec 2000, PENDING

DATE NUMBER \_\_\_\_\_

US 1999-169629P 19991208 (60) PRIORITY INFORMATION: US 2000-187470P 20000306 (60)

DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT:

Frank C. Eisenschenk, Ph.D., SALIWANCHIK, LLOYD & LEGAL REPRESENTATIVE:

SALIWANCHIK, 2421 N.W. 41 STREET, SUITE A-1,

GAINESVILLE, FL, 32606-6669

22 NUMBER OF CLAIMS: 1 EXEMPLARY CLAIM:

EXEMPLARY CLAIM:

NUMBER OF DRAWINGS:

TIME COUNT:

- 27600

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The invention concerns GENSET polynucleotides and polypeptides. Such GENSET products may be used as reagents in forensic analyses, as chromosome markers, as tissue/cell/organelle-specific markers, in the production of expression vectors. In addition, they may be used in screening and diagnosis assays for abnormal GENSET expression and/or biological activity and for screening compounds that may be used in the treatment of GENSET-related disorders.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 16 OF 22 USPATFULE on STN

2003:38352 USPATFULL ACCESSION NUMBER:

TITLE: 143 human secreted proteins

Rosen, Craig A., Laytonsville, MD, UNITED STATES INVENTOR(S):

Ruben, Steven M., Olney, MD, UNITED STATES Moore, Paul A., Germantown, MD, UNITED STATES Young, Paul E., Gaithersburg, MD, UNITED STATES Komatsoulis, George A., Silver Spring, MD, UNITED

STATES

Birse, Charles E., North Potomac, MD, UNITED STATES

Duan, Roxanne D., Bethesda, MD, UNITED STATES

Florence, Kimberly A., Rockville, MD, UNITED STATES Soppet, Daniel R., Centreville, VA, UNITED STATES

NUMBER KIND DATE US 2003027999 A1 20030206 US 2001-986480 A1 20011108 (9)

APPLICATION INFO.: Continuation-in-part of Ser. No. WO 2000-US12788, RELATED APPLN. INFO.:

filed on 11 May 2000, UNKNOWN

NUMBER DATE

PRIORITY INFORMATION: US 1999-134068P 19990513 (60) Utility

DOCUMENT TYPE:

FILE SEGMENT: APPLICATION
LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 24 EXEMPLARY CLAIM: 1 LINE COUNT: 29687

PATENT INFORMATION:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to novel human secreted proteins and isolated nucleic acids containing the coding regions of the genes encoding such proteins. Also provided are vectors, host cells, antibodies, and recombinant methods for producing human secreted proteins. The invention further relates to diagnostic and therapeutic methods useful for diagnosing and treating diseases, disorders, and/or conditions related to these novel human secreted proteins.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 17 OF 22 USPATFULL on STN

ACCESSION NUMBER: 2002:192264 USPATFULL

TITLE: Staphylococcus aureus polynucleotides and

polypeptides

INVENTOR(S): Choi, Gil H., Rockville, MD, UNITED STATES

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. WO 2000-US23773,

filed on 31 Aug 2000, UNKNOWN Continuation-in-part of Ser. No. US 1997-781986, filed on 3 Jan 1997, PENDING Continuation-in-part of Ser. No. US

1997-956171, filed on 20 Oct 1997, PENDING
NUMBER DATE

PRIORITY INFORMATION: US 1999-151933P 19990901 (60)
US 1996-9861P 19960105 (60)
US 1996-9861P 19960105 (60)

DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 96
EXEMPLARY CLAIM: 1
LINE COUNT: 9945

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to novel genes from S. aureus and the polypeptides they encode. Also provided are vectors, host cells, antibodies and recombinant methods for producing the same. The invention further relates to screening methods for identifying agonists and antagonists of S. aureus polypeptide activity. The invention additionally relates to diagnostic methods for detecting Staphylococcus nucleic acids, polypeptides and antibodies in a biological sample. The present invention further relates to novel vaccines for the prevention or attenuation of infection by Staphylococcus.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 18 OF 22 USPATFULL on STN

ACCESSION NUMBER: 2002:141608 USPATFULL

TITLE: Nucleotide sequence of Escherichia coli

pathogenicity islands

INVENTOR(S): Dillon, Patrick J., Carlsbad, CA, UNITED STATES

Choi, Gil H., Rockville, MD, UNITED STATES

Welch, Rodney A., Madison, WI, UNITED STATES

PATENT ASSIGNEE(S): Human Genome Sciences, Inc., Rockville, MD, UNITED

STATES (U.S. corporation)

RELATED APPLN. INFO.: Division of Ser. No. US 1997-976259, filed on 21

Nov 1997, GRANTED, Pat. No. US 6316609

NUMBER DATE \_\_\_\_\_

PRIORITY INFORMATION: US 1997-61953P 19971014 (60) US 1996-31626P 19961122 (60)

DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT:

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1

2 Drawing Page(s) 8481 NUMBER OF DRAWINGS:

LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to novel genes located in two chromosomal regions within uropathogenic E. coli that are associated with virulence. These chromosomal regions are known as pathogenicity islands (PAIs). In particular, the present application discloses 142 sequenced fragments (contigs) of DNA from two pools of cosmids covering pathogenicity islands PAI IV and PAI V located on the chromosome of the uropathogenic Escherichia coli J96. Further disclosed are 351 predicted protein-coding open reading frames within the sequenced fragments.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 19 OF 22 USPATFULL on STN

ACCESSION NUMBER: 2002:140861 USPATFULL

Soluble CD1 compositions and uses thereof TITLE:

Gumperz, Jenny E., Jamaica Plain, MA, UNITED STATES INVENTOR(S):

Brenner, Michael B., Newton, MA, UNITED STATES Behar, Samuel M., Needham, MA, UNITED STATES

NUMBER KIND DATE US 2002071842 A1 20020613 US 2001-874470 A1 20010605 (9) PATENT INFORMATION: APPLICATION INFO.:

NUMBER DATE

\_\_\_\_\_ PRIORITY INFORMATION: US 2000-209416P 20000605 (60)

DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT:

LEGAL REPRESENTATIVE: Elizabeth R. Plumer, c/o Wolf, Greenfield & Sacks,

P.C., Federal Reserve Plaza, 600 Atlantic Avenue,

Boston, MA, 02210-2211

66 . NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1 2798 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Compositions and methods for identifying CD1 antigens and AΒ CD1-restricted T cells, and diagnostic and therapeutic uses of same are provided. The compositions include CD1 fusion proteins, preferably multivalent fusion proteins that are present in multimeric form (e.g., by Protein A binding multiple CD1 fusion

proteins).

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 20 OF 22 USPATFULL on STN ACCESSION NUMBER: 2002:19393 USPATFULL Secreted protein HLHFP03 TITLE: Rosen, Craig A., Laytonsville, MD, United States INVENTOR(S): Ruben, Steven M., Olney, MD, United States Olsen, Henrik S., Gaithersburg, MD, United States Ebner, Reinhard, Gaithersburg, MD, United States PATENT ASSIGNEE(S): Human Genome Sciences, Inc., Rockville, MD, United States (U.S. corporation) NUMBER KIND DATE \_\_\_\_\_ US 6342581 B1 US 1999-227357 20020129 PATENT INFORMATION: 19990108 (9) APPLICATION INFO.: Continuation-in-part of Ser. No. WO 1998-US13684, RELATED APPLN. INFO.: filed on 7 Jul 1998 DATE NUMBER US 1997-58785P 19970912 (60) PRIORITY INFORMATION: 19970912 (60) US 1997-58664P 19970912 (60) US 1997-58660P US 1997-58661P 19970912 (60) US 1997-55722P 19970818 (60) US 1997-55723P 19970818 (60) US 1997-55948P 19970818 (60) 19970818 (60) US 1997-55949P US 1997-55953P 19970818 (60) 19970818 (60) US 1997-55950P US 1997-55947P 19970818 (60) US 1997-55964P 19970818 (60) US 1997-56360P 19970818 (60) US 1997-55684P 19970818 (60) US 1997-55984P 19970818 (60) US 1997-55954P 19970818 (60) US 1997-51926P 19970708 (60) US 1997-52793P 19970708 (60) US 1997-51925P 19970708 (60) 19970708 (60) US 1997-51929P US 1997-52803P 19970708 (60) 19970708 (60) US 1997-52732P US 1997-51931P 19970708 (60) US 1997-51932P 19970708 (60) - US 1997-51916P 19970708 (60) 19970708 (60) US 1997-51930P 19970708 (60) US 1997-51918P US 1997-51920P 19970708 (60) US 1997-52733P 19970708 (60) US 1997-52795P 19970708 (60) US 1997-51919P 19970708 (60) US 1997-51928P 19970708 (60) DOCUMENT TYPE: Utility FILE SEGMENT: GRANTED Myers, Carla J. PRIMARY EXAMINER: Spiegler, Alexander H. ASSISTANT EXAMINER: Human Genome Sciences, Inc. LEGAL REPRESENTATIVE: NUMBER OF CLAIMS: 46 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 0 Drawing Figure(s); 0 Drawing Page(s)

LINE COUNT: 18742

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to novel human secreted proteins and isolated nucleic acids containing the coding regions of the genes encoding such proteins. Also provided are vectors, host cells, antibodies, and recombinant methods for producing human secreted proteins. The invention further relates to diagnostic and therapeutic methods useful for diagnosing and treating disorders related to these novel human secreted proteins.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 21 OF 22 USPATFULL on STN

ACCESSION NUMBER: 96:50802 USPATFULL

TITLE: Cytolysin gene and gene product

INVENTOR(S): Goebel, Werner, Veitschochheim, Germany, Federal

Republic of

Libby, Stephen J., San Diego, CA, United States

Heffron, Fred, Portland, OR, United States

PATENT ASSIGNEE(S): Merck Patent Gesellschaft mit beschrankter Haftung,

Darmstadt, Germany, Federal Republic of (non-U.S.

corporation)

DOCUMENT TYPE: Utility
FILE SEGMENT: Granted

PRIMARY EXAMINER: Vogel, Nancy T.

LEGAL REPRESENTATIVE: Millen, White, Zelano, & Branigan

NUMBER OF CLAIMS: 6
EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 6 Drawing Figure(s); 5 Drawing Page(s)

LINE COUNT: 1378

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AS almonella gene, encoding a cytolysin, has been identified by screening for hemolysis on blood agar. The gene (slyA) is present in every strain of Salmonella examined in Shigella, and enteroinvasive Escherichia coli (EIEC) but not in other enterobacteriaceae. It is encoded near 28.5 minutes on the chromosome. A SlyA (salmolysin) has hemolytic and cytolytic activity and has a molecular weight predicted by the DNA sequence. LD.sub.50 and infection kinetics data in mice indicate that the toxin is required for virulence and facilitates Salmonella survival within peritoneal macrophages.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 22 OF 22 USPATFULL on STN

ACCESSION NUMBER: 94:44553 USPATFULL

TITLE: Process for converting lipid-containing bacterial

capsular polysaccharide into lipid-free

polysaccharide

INVENTOR(S): Lee, Ann L., Lansdale, PA, United States

Rienstra, Mark S., Lansdale, PA, United States Manger, Walter E., Harleysville, PA, United States

Sitrin, Robert D., Lafayette Hill, PA, United

States

Merck & Co., Inc., Rahway, NJ, United States (U.S. PATENT ASSIGNEE(S):

corporation)

NUMBER KIND DATE -----US 5314811 PATENT INFORMATION: 19940524 US 1992-909346 19920713 (7) APPLICATION INFO.:

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

Griffin, Ronald W. PRIMARY EXAMINER:

Bencen, Gerard H., Tribble, Jack L., Matukaitis, LEGAL REPRESENTATIVE:

Paul D.

NUMBER OF CLAIMS: 14 EXEMPLARY CLAIM:

3 Drawing Figure(s); 3 Drawing Page(s) NUMBER OF DRAWINGS:

LINE COUNT: 1440

CAS INDEXING IS AVAILABLE 'FOR THIS PATENT.

A process for converting lipid-containing bacterial capsular polysaccharide, such as lipo-polyribosyl ribitol phosphate, lipo-PRP, into lipid-free, endotoxin-free polysaccharide, such as polyribosyl ribitol phosphate, PRP, by solubilizing polysaccharide-containing powder derived from culture media of bacteria, such as Haemophilus influenzae type b, cleaving covalently bound fatty acids from the polysaccharide, and removing the lipids, and endotoxin.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

(FILE 'CAPLUS, MEDLINE, BIOSIS, EMBASE, WPIDS, CONFSCI, SCISEARCH, JICST-EPLUS, JAPIO, USPATFULL' ENTERED AT 10:52:44 ON 03 MAY 2006)

1248 S "APICELLA M"?/AU L22 Author (5) 19139 S "EDWARDS J"?/AU L23

62 S L22 AND L23 L24

L25 13 S (L22 OR L23 OR L24) AND (L2 OR PLD) L26 8 DUP REM L25 (5 DUPLICATES REMOVED)

L26 ANSWER 1 OF 8 WPIDS COPYRIGHT 2006 THE THOMSON CORP on STN

ACCESSION NUMBER:

2005-123122 [13] WPIDS

CROSS REFERENCE:

2002-619227 [66]

DOC. NO. CPI:

C2005-040896

TITLE:

New transgenic Neisseria bacterium comprising a

disrupted pld gene and a reduced phospholipase D activity, useful

for preventing or treating neisserial infections,

such as gonorrhea.

DERWENT CLASS:

B04 D16

INVENTOR(S): PATENT ASSIGNEE(S): \_APICELLA, M A; EDWARDS, J L (IOWA) UNIV IOWA RES FOUND

COUNTRY COUNT: 107

PATENT INFORMATION:

KIND DATE PATENT NO WEEK LA PG \_\_\_\_\_\_

WO 2005010036 A1 20050203 (200513)\* EN 163

RW: AT BE BG BW CH CY CZ DE DK EA EE ES FI FR GB GH GM GR HU IE IT KE LS LU MC MW MZ NA NL OA PL PT RO SD SE SI SK SL SZ TR TZ UG

W: AE AG AL AM AT AU AZ BA BB BG BR BW BY BZ CA CH CN CO CR CU CZ DE DK DM DZ EC EE EG ES FI GB GD GE GH GM HR HU ID IL IN IS JP

KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NA NI NO NZ OM PG PH PL PT RO RU SC SD SE SG SK SL SY TJ TM TN TR TT TZ UA UG UZ VC VN YU ZA ZM ZW

#### APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
WO 2005010036	A1	WO 2004-US22708	20040715

PRIORITY APPLN. INFO: US 2003-665990 20030919; US

2003-621184 20030715

AN 2005-123122 [13] WPIDS

CR 2002-619227 [66]

AB W02005010036 A UPAB: 20050224

NOVELTY - A transgenic Neisseria bacterium comprising a disrupted pld gene, is new. The bacterium has reduced phospholipase D (PLD) activity as compared

to the **phospholipase D** activity of a corresponding wild-type Neisseria.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated and purified polynucleotide encoding a PLD from a Neisseria bacterium;
- (2) an isolated and purified polypeptide that is encoded by the above polynucleotide and that comprises **phospholipase**D from a Neisseria bacterium;
- (3) a vaccine comprising an immunogenic amount of a **PLD** polypeptide from Neisseria, which amount immunizes a patient against a neisserial infection, in combination with a physiological, non-toxic vehicle;
- (4) protecting a patient against Neisseria colonization or infection, comprising administering to the patient an amount of the vaccine mentioned above; and
- (5) preventing infection or colonization of Neisseria in a patient by administering to the patient a compound that inhibits neisserial phospholipase D.

ACTIVITY - Antibacterial; Gynecological.

No biological data given.

MECHANISM OF ACTION - Vaccine.

USE - The composition and methods are useful for preventing or treating neisserial infections, such as gonorrhea. Dwg.0/23

L26 ANSWER 2 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 1

ACCESSION NUMBER: 2004:1080507 CAPLUS

DOCUMENT NUMBER: 142:54745

TITLE: . Vaccine and compositions comprising a neisserial

phospholipase D for the

prevention and treatment of neisserial infections

INVENTOR(S): Apicella, Michael A.; Edwards,

Jennifer L.

PATENT ASSIGNEE(S): USA

SOURCE: U.S. Pat. Appl. Publ., 103 pp., Cont.-in-part of

U.S. Ser. No. 621,184.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2 PATENT INFORMATION:

PA.	PATENT NO.		KIND DATE		APPLICATION NO.						DATE					
US	2003	1000	71		A1		20030529		US 2003-665990 US 2002-66551 WO 2004-US22708			1	20020131			
WO		AE, CH, GB, KR,	AG, CN, GD, KZ,	AL, CO, GE, LC,	AM, CR, GH, LK,	AT, CU, GM, LR,	AU, CZ, HR, LS,	AZ, DE, HU, LT,	BA, DK, ID, LU,	BB, DM, IL, LV,	BG, DZ, IN, MA,	BR, EC, IS, MD,	BW, EE, JP, MG,	BY, EG, KE, MK,	BZ, ES, KG, MN,	CA, FI, KP, MW,
	DW.	SE, VN,	SG, YU,	SK, ZA,	SL, ZM,	SY, ZW	NZ, TJ, MW,	TM,	TN,	TR,	TT,	TZ,	UA,	υG,	UZ,	VC,
	100.	AM, DE, PT,	AZ, DK, RO,	BY, EE, SE,	KG, ES, SI,	KZ, FI, SK,	MD, FR, TR,	RU, GB, BF,	TJ, GR,	TM, HU,	AT, IE,	BE, IT,	BG, LU,	CH, MC,	CY, NL,	CZ, PL,
PRIORITY	Y APP	•	•	•	,	J.,,	12,		1	US 2	001-2	2660'	70P	;	P 2	0010131
									Ţ	US 2	001-	3103	56P	:	P 2	0010806
				_					1	US 2	001-	3444	52P		P 2	0011023
									1	US 2	002-	6655	1	1	A2 2	0020131
									1	US 2	003-	6211	8 4	i	A2 2	0030715
									1	US 2	003-	6659	90	j	A2 2	0030919

The present invention provides a polypeptide, polynucleotide, vaccine, and a method of vaccination effective to immunize a mammal against a neisserial infection, e.g., an infection caused by Neisseria gonorrhoeae or Neisseria meningitidis by using a neisserial phospholipase D (PLD) polypeptide in combination with a physiol.—acceptable, non-toxic vehicle. In addition, the invention provides a transgenic Neisseria bacterium comprising a disrupted pld gene wherein the bacterium has reduced phospholipase D activity as compared to the phospholipase D activity of a corresponding wild-type Neisseria.

L26 ANSWER 3 OF 8 USPATFULL on STN

ACCESSION NUMBER:

2004:139122 USPATFULL

TITLE:

Method of removing silicon oxide from a surface of

a substrate

INVENTOR(S):

Hu, Xiaoming, Chandler, AZ, UNITED STATES Craigo, James B., Tempe, AZ, UNITED STATES

Droopad, Ravindranath, Chandler, AZ, UNITED STATES

Edwards, John L., JR., Phoenix, AZ,

UNITED STATES

Liang, Yong, Gilbert, AZ, UNITED STATES Wei, Yi, Chandler, AZ, UNITED STATES Yu, Zhiyi, Gilbert, AZ, UNITED STATES

NUMBER	KIND	DATE

PATENT INFORMATION: US 2004106296 A1 20040603

US 6806202 B2 20041019

APPLICATION INFO.: US 2002-309500 A1 20021203 (10)

DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: OBLON, SPIVAK, MCCLELLAND, MAIER & NEUSTADT, P.C.,

1940 DUKE STREET, ALEXANDRIA, VA, 22314

NUMBER OF CLAIMS: 20 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 1 Drawing Page(s)

LINE COUNT: 331

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A method for removing silicon oxide from a surface of a substrate is disclosed. The method includes depositing material onto the silicon oxide (110) and heating the substrate surface to a sufficient temperature to form volatile compounds including the silicon oxide and the deposited material (120). The method also includes heating the surface to a sufficient temperature to remove any remaining deposited material (130).

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L26 ANSWER 4 OF 8 CAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 2

ACCESSION NUMBER: 2003:873418 CAPLUS

DOCUMENT NUMBER: 139:379737

TITLE: Gonococcal phospholipase D

modulates the expression and function of complement receptor 3 in primary cervical

epithelial cells

AUTHOR(S): Edwards, Jennifer L.; Entz, David D.;

Apicella, Michael A.

CORPORATE SOURCE: Department of Microbiology, University of Iowa,

Iowa City, IA, 52242, USA

SOURCE: Infection and Immunity (2003), 71(11), 6381-6391

CODEN: INFIBR; ISSN: 0019-9567

PUBLISHER: American Society for Microbiology

DOCUMENT TYPE: Journal LANGUAGE: - English

AB CR3-mediated endocytosis is a primary mechanism by which Neisseria gonorrhoeae elicits membrane ruffling and cellular invasion of the cervical epithelia. The authors' data indicate that, upon infection of cervical epithelia, N. gonorrhoeae specifically releases proteins,

including a phospholipase D (PLD) homolog, which facilitate membrane ruffling. To elucidate the function of gonococcal PLD in infection of the cervical epithelia, the authors constructed an N. gonorrhoeae PLD mutant. By comparative association and/or invasion assays, the authors demonstrated that PLD mutant gonococci are impaired in their ability to adhere to and to invade primary cervical cells. This defect can be rescued by the addition of supernatants obtained from wild-type-infected cell monolayers but not by exogenously added Streptomyces PLD. The decreased level of total cell association (i.e., adherence and invasion) observed for mutant gonococci is, in part, attributed to the inability of these bacteria to recruit CR3 to the cervical cell surface with extended infection. Using electron microscopy, the authors demonstrate that gonococcal PLD may be necessary to potentiate membrane ruffling and clustering of gonococci on the cervical cell surface. These data may be indicative of the inability of PLD mutant gonococci to recruit CR3 to

the cervical cell surface. Alternatively, in the absence of gonococcal PLD, signal transduction events required for CR3 clustering may not be activated. Collectively, the authors' data indicate that PLD augments CR3-mediated gonococcus invasion of and survival within cervical epithelia.

REFERENCE COUNT:

THERE ARE 33 CITED REFERENCES AVAILABLE FOR 33

THIS RECORD. ALL CITATIONS AVAILABLE IN THE

RE FORMAT

L26 ANSWER 5 OF 8 USPATFULL on STN

ACCESSION NUMBER:

2002:215158 USPATFULL

TITLE:

Method and apparatus for efficiently moving

portions of a memory block

INVENTOR(S):

Somers, Jeffrey, Northboro, MA, UNITED STATES Alden, Andrew, Leominster, MA, UNITED STATES Edwards, John, Clinton, MA, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 2002116555	A1	20020822	
APPLICATION INFO.:	US 6948010 US 2000-742989	B2 A1	20050920 20001220	(9)
DOCUMENT TYPE:	Utility			, - ,
FILE SEGMENT:	APPLICATION			

LEGAL REPRESENTATIVE: TESTA, HURWITZ & THIBEAULT, LLP, HIGH STREET TOWER,

125 HIGH STREET, BOSTON, MA, 02110 23

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS:

6 Drawing Page(s)

LINE COUNT: 687

AB

The present invention relates to a method and system for transferring portions of a memory block. A first data mover is configured with a first start address corresponding to a first portion of a source memory block. A second data mover is configured with a second start address corresponding to a second portion of the source memory block sized differently from the first portion. The first portion of the source memory block is transferred by the first data mover and the second portion of the source memory block is transferred by the second data mover.

L26 ANSWER 6 OF 8 USPATFULL on STN

ACCESSION NUMBER:

2002:170289 USPATFULL

TITLE:

Low leakage current metal oxide-nitrides and method

of fabricating same

INVENTOR(S):

Yu, Zhiyi, Gilbert, AZ, UNITED STATES

Droopad, Ravindranath, Chandler, AZ, UNITED STATES

Overgaard, Corey, Phoenix, AZ, UNITED STATES Edwards, John Leonard, JR., Phoenix, AZ,

UNITED STATES

PATENT ASSIGNEE(S):

Motorola, Inc. (U.S. corporation)

	NUMBER	KIND	DATE	
PATENT INFORMATION: APPLICATION INFO.: DOCUMENT TYPE:	US 2002089023 US 2001-755691 Utility	A1 A1	20020711 20010105	(9)

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

OBLON SPIVAK MCCLELLAND MAIER & NEUSTADT PC, FOURTH

FLOOR, 1755 JEFFERSON DAVIS HIGHWAY, ARLINGTON, VA,

22202

83 NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 1 Drawing Page(s)

LINE COUNT: 1033

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A structure and method for forming a high dielectric constant device structure includes a monocrystalline semiconductor substrate and an insulating layer formed of a metal oxide-nitride such as M.sub.nO.sub.m-xN.sub.x, wherein M is a metallic or semi-metallic element or combination of metallic and/or semi-metallic elements and m and n are integers. Semiconductor devices formed in accordance with the present invention exhibit low leakage current density and improved chemical, thermal, and electrical stability over conventional metal oxides.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L26 ANSWER 7 OF 8 USPATFULL on STN

ACCESSION NUMBER: 2002:141270 USPATFULL

Method of removing an amorphous oxide from a TITLE:

monocrystalline surface

Edwards, John L., JR., Phoenix, AZ, INVENTOR(S):

UNITED STATES

Wei, Yi, Chandler, AZ, UNITED STATES

Jordan, Dirk C., Gilbert, AZ, UNITED STATES Hu, Xiaoming, Chandler, AZ, UNITED STATES

Craigo, James Bradley, Tempe, AZ, UNITED STATES Droopad, Ravindranath, Chandler, AZ, UNITED STATES

Yu, Zhiyi, Gilbert, AZ, UNITED STATES

Demkov, Alexander A., Phoenix, AZ, UNITED STATES MOTOROLA, INC., Schaumburg, IL, 60196-1079 (U.S.

PATENT ASSIGNEE(S): corporation)

KIND NUMBER DATE US 2002072253 A1 20020613 PATENT INFORMATION: US 6693033 B2 20040217 APPLICATION INFO.: - US 2001-983854 A1 20011026 (9)

Continuation-in-part of Ser. No. US 2000-502023, RELATED APPLN. INFO.:

filed on 10 Feb 2000, PENDING

DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT:

OBLON SPIVAK MCCLELLAND MAIER & NEUSTADT PC, FOURTH LEGAL REPRESENTATIVE:

FLOOR, 1755 JEFFERSON DAVIS HIGHWAY, ARLINGTON, VA,

22202

NUMBER OF CLAIMS: 29 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 1 Drawing Page(s)

LINE COUNT: 448

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A method of removing an amorphous oxide from a surface of a monocrystalline substrate is provided. The method includes depositing a passivation material overlying the amorphous oxide. The monocrystalline substrate is then heated so that the amorphous oxide layer decomposes into at least one volatile species that is liberated from the surface.

> : Shears 571-272-2528 Searcher

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L26 ANSWER 8 OF 8 SCISEARCH COPYRIGHT (c) 2006 The Thomson Corporation

on STN

ACCESSION NUMBER: 1995:384846 SCISEARCH

THE GENUINE ARTICLE: RB325

TITLE: ACCUMULATION OF PHOSPHATIDYLALCOHOL IN CULTURED-CELLS

- USE OF SUBCELLULAR FRACTIONATION TO INVESTIGATE

PHOSPHOLIPASE-D ACTIVITY DURING

SIGNAL-TRANSDUCTION

AUTHOR: EDWARDS J S (Reprint); MURRAY A W

CORPORATE SOURCE: FLINDERS UNIV S AUSTRALIA, SCH BIOL SCI, POB 2100,

ADELAIDE, SA 5001, AUSTRALIA (Reprint)

COUNTRY OF AUTHOR: AUSTRALIA

SOURCE: BIOCHEMICAL JOURNAL, (1 JUN 1995) Vol. 308, Part 2,

pp. 473-480. ISSN: 0264-6021.

PUBLISHER: PORTLAND PRESS, 59 PORTLAND PLACE, LONDON W1N 3AJ,

ENGLAND.

DOCUMENT TYPE: Article; Journal

LANGUAGE: English

REFERENCE COUNT: 44

ENTRY DATE: Entered STN: 1995

Last Updated on STN: 1995

\*ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS\*

AB Phosphatidylalcohol accumulates as a product of a

phospholipase D (PLD)-catalysed transphosphatidylation reaction in cells incubated in the presence of a primary alcohol, In the presence of ethanol the phorbol ester, phorbol 12-myristate 13-acetate (PMA), stimulated the accumulation of [H-3] phosphatidylethanol (PEth) in HeLa cells prelabelled with [H-3] palmitic acid. Radioactivity associated with PEth increased linearly during a 30 min incubation, indicating that a sustained activation of PLD is caused by PMA in these cells. This was accompanied by the membrane association of protein kinase C-alpha (PKC-alpha), the PKC isoform that recent studies indicate is involved in the activation of PLD. In similar experiments, the neuropeptide bradykinin stimulated an accumulation of PEth in 3T3 Li cells. The radioactivity associated with PEth increased to a maximal level at 30 s and plateaued after this time, suggesting that bradykinin induces only a transient activation of PLD in these cells. This is consistent with the effects of bradykinin on PKC-alpha, which underwent a rapid and transient association with cell membranes. The subcellular localization of PEth was examined using the technique of subcellular fractionation on Percoll density gradients to isolate organelle-enriched fractions from HeLa and 3T3 Li cells. An accumulation of [H-3]PEth was measured in the plasma-membrane (PM)-enriched fractions of both HeLa and 3T3 Li cells after incubation with PMA and bradykinin respectively. This was accompanied by a time-dependent accumulation of [H-3] PEth in the combined mitochondrial and endoplasmic reticulum (MER)-enriched fractions of both cell lines. PMA was also found to cause translocation of PKC-alpha to both the PM- and MER-enriched fractions in HeLa cells. However, bradykinin stimulated the translocation of PKC-alpha to the PM-enriched fractions only of 3T3 Li cells. results show that PLD activation leads to the accumulation of PEth in both the PM and MER fractions. We therefore propose that either bradykinin activates a PM-associated PLD and the PLD reaction product is rapidly translocated to other membrane

systems or it activates an MER-associated  ${\tt PLD}$  by a mechanism that does not involve PKC-alpha.

FILE 'HOME' ENTERED AT 10:54:22 ON 03 MAY 2006

### => d his ful

L7

(FILE 'HOME' ENTERED AT 10:43:42 ON 03 MAY 2006) SET COST OFF

FILE 'REGISTRY' ENTERED AT 10:43:53 ON 03 MAY 2006 E PHOSPHOLIPASE D/CN 5

. L1 154 SEA ABB=ON PLU=ON PHOSPHOLIPASE D ?/CN

FILE 'CAPLUS' ENTERED AT 10:44:53 ON 03 MAY 2006

L2 4852 SEA ABB=ON PLU=ON L1 OR (PHOSPHOLIPASE OR PHOSPHO LIPASE OR LECITHINASE) (1W) D OR (PHOSPHATIDYLCHOLINE OR PHOSPHATIDY L CHOLINE) (W) (PHOSPHOHYDROLASE OR PHOSPHO HYDROLASE)

L3 8 SEA ABB=ON PLU=ON L2 AND NEISSER? L4 8 SEA ABB=ON PLU=ON L2 AND ?NEISSER?

FILE 'REGISTRY' ENTERED AT 10:46:09 ON 03 MAY 2006

FILE 'CAPLUS' ENTERED AT 10:46:09 ON 03 MAY 2006
D QUE L4
D L4 1-8 .BEVSTR

FILE 'MEDLINE, BIOSIS, EMBASE, WPIDS, CONFSCI, SCISEARCH, JICST-EPLUS, JAPIO' ENTERED AT 10:46:11 ON 03 MAY 2006

L5 11 SEA ABB=ON PLU=ON L4

L6 5 DUP REM L5 (6 DUPLICATES REMOVED)
D 1-5 IBIB ABS

FILE 'CAPLUS' ENTERED AT 10:47:01 ON 03 MAY 2006

2 SEA ABB=ON PLU=ON PLD AND NEISSER?

L8 0 SEA ABB=ON PLU=ON L7 NOT L4

FILE 'MEDLINE, BIOSIS, EMBASE, WPIDS, CONFSCI, SCISEARCH, JICST-EPLUS, JAPIO' ENTERED AT 10:47:23 ON 03 MAY 2006

L9 9 SEA ABB=ON PLU=ON L7

L10 4 SEA ABB=ON PLU=ON L9 NOT L5

L11 4 DUP REM L10 (0 DUPLICATES REMOVED)

D KWIC

D KWIC 2-3

L12 1 SEA ABB=ON PLU=ON L11 AND (POLYPEPTIDE OR PEPTIDE OR PROTEIN OR POLYPROTEIN)

D KWIC

L13 0 SEA ABB=ON PLU=ON L12 AND (VACCIN? OR IMMUNIS? OR IMMUNIZ?)

FILE 'MEDLINE' ENTERED AT 10:49:32 ON 03 MAY 2006

L14 0 SEA ABB=ON PLU=ON (PHOSPHOLIPASE D AND NEISSERIA)/CT D QUE

L15 6 SEA ABB=ON PLU=ON (PHOSPHOLIPASE D AND BACTERIA)/CT D QUE D 1-6 .BEVERLYMED

FILE 'USPATFULL' ENTERED AT 10:50:39 ON 03 MAY 2006

L16 39 SEA ABB=ON PLU=ON (L2 OR PLD) (L) NEISSER?

L17 39 SEA ABB=ON PLU=ON L16(L)(POLYPEPTIDE OR PEPTIDE OR PROTEIN OR POLYPROTEIN)

L18 27 SEA ABB=ON PLU=ON L17(L) (VACCIN? OR IMMUNIS? OR IMMUNIZ?)

L19 564 SEA ABB=ON PLU=ON (L2 OR PLD)(S)(POLYPEPTIDE OR PEPTIDE

L20 L21	OR PROTEIN OR POLYPF 23 SEA ABB=ON PLU=ON 22 SEA ABB=ON PLU=ON	,
	D OUE	
	D 1-22 IBIB ABS	
	FILE 'CAPLUS, MEDLINE, BIOSIS,	EMBASE, WPIDS, CONFSCI, SCISEARCH,
	JICST-EPLUS, JAPIO, USPATFULL'	ENTERED AT 10:52:44 ON 03 MAY 2006
L22	1248 SEA ABB=ON PLU=ON	"APICELLA M"?/AU
L23	19139 SEA ABB=ON PLU=ON	"EDWARDS J"?/AU
L24	62 SEA ABB=ON PLU=ON	L22 AND L23
L25	13 SEA ABB=ON PLU=ON	(L22 OR L23 OR L24) AND (L2 OR PLD)
L26	8 DUP REM L25 (5 DUPLI	CATES REMOVED)
	D 1-8 IBIB ABS	

FILE 'HOME' ENTERED AT 10:54:22 ON 03 MAY 2006

FILE HOME

# FILE REGISTRY

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 2 MAY 2006 HIGHEST RN 882569-16-6 DICTIONARY FILE UPDATES: 2 MAY 2006 HIGHEST RN 882569-16-6

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### FILE CAPLUS

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FILE COVERS 1907 - 3 May 2006 VOL 144 ISS 19 FILE LAST UPDATED: 2 May 2006 (20060502/ED)

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### FILE MEDLINE

FILE LAST UPDATED: 2 MAY 2006 (20060502/UP). FILE COVERS 1950 TO DAT

On December 11, 2005, the 2006 MeSH terms were loaded.

The MEDLINE reload for 2006 is now (26 Feb.) available. For details on the 2006 reload, enter HELP RLOAD at an arrow prompt (=>). See also:

http://www.nlm.nih.gov/mesh/

http://www.nlm.nih.gov/pubs/techbull/nd04/nd04\_mesh.html

http://www.nlm.nih.gov/pubs/techbull/nd05/nd05\_med\_data\_changes.ht

http://www.nlm.nih.gov/pubs/techbull/nd05/nd05\_2006\_MeSH.html

OLDMEDLINE is covered back to 1950.

MEDLINE thesauri in the /CN, /CT, and /MN fields incorporate the MeSH 2006 vocabulary.

This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE BIOSIS

FILE COVERS 1969 TO DATE.

CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNs) PRESENT FROM JANUARY 1969 TO DATE.

RECORDS LAST ADDED: 26 April 2006 (20060426/ED)

## FILE EMBASE

FILE COVERS 1974 TO 2 May 2006 (20060502/ED)

EMBASE has been reloaded. Enter HELP RLOAD for details.

EMBASE is now updated daily. SDI frequency remains weekly (default) and biweekly.

This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE WPIDS

FILE LAST UPDATED: 2 MAY 2006 <20060502/UP>
MOST RECENT DERWENT UPDATE: 200628 <200628/DW>

DERWENT WORLD PATENTS INDEX SUBSCRIBER FILE, COVERS 1963 TO DATE

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FILE CONFSCI

FILE COVERS 1973 TO 10 Apr 2006 (20060410/ED)

CSA has resumed updates, see NEWS FILE

FILE SCISEARCH

FILE COVERS 1974 TO 28 Apr 2006 (20060428/ED)

SCISEARCH has been reloaded, see HELP RLOAD for details.

FILE JICST-EPLUS

FILE COVERS 1985 TO 1 MAY 2006 (20060501/ED)

THE JICST-EPLUS FILE HAS BEEN RELOADED TO REFLECT THE 1999 CONTROLLED TERM (/CT) THESAURUS RELOAD.

FILE JAPIO

FILE LAST UPDATED: 3 APR 2006 <20060403/UP>
FILE COVERS APRIL 1973 TO DECEMBER 22, 2005

- >>> GRAPHIC IMAGES AVAILABLE <<<
- >>> NEW IPC8 DATA AND FUNCTIONALITY NOT YET AVAILABLE IN THIS FILE.

  USE IPC7 FORMAT FOR SEARCHING THE IPC. WATCH THIS SPACE FOR FURTHE

  DEVELOPMENTS AND SEE OUR NEWS SECTION FOR FURTHER INFORMATION

  ABOUT THE IPC REFORM <<<

FILE USPATFULL

FILE COVERS 1971 TO PATENT PUBLICATION DATE: 2 May 2006 (20060502/PD)

FILE LAST UPDATED: 2 May 2006 (20060502/ED)

HIGHEST GRANTED PATENT NUMBER: US7039955

HIGHEST APPLICATION PUBLICATION NUMBER: US2006090232

CA INDEXING IS CURRENT THROUGH 2 May 2006 (20060502/UPCA)

ISSUE CLASS FIELDS (/INCL) CURRENT THROUGH: 2 May 2006 (20060502/PD)

REVISED CLASS FIELDS (/NCL) LAST RELOADED: Feb 2006

USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Feb 2006